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(54) Title: NOVEL NUCLEIC ACIDS AND POLYPEPTIDES

• (57) Abstract: The present invention provides novel nucleic acids, novel polypeptide sequences encoded by these nucleic acids and uses thereof.



NOVEL NUCLEIC ACIDS AND POLYPEPTIDES

1. TECHNICAL FIELD

The present invention provides novel polynucleotides and proteins encoded by such polynucleotides, along with uses for these polynucleotides and proteins, for example in therapeutic, diagnostic and research methods.

2. BACKGROUND

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Technology aimed at the discovery of protein factors (including e.g., cytokines, such as lymphokines, interferons, CSFs, chemokines, and interleukins) has matured rapidly over the past decade. The now routine hybridization cloning and expression cloning techniques clone novel polynucleotides "directly" in the sense that they rely on information directly related to the discovered protein (i.e., partial DNA/amino acid sequence of the protein in the case of hybridization cloning; activity of the protein in the case of expression cloning). More recent "indirect" cloning techniques such as signal sequence cloning, which isolates DNA sequences based on the presence of a now well-recognized secretory leader sequence motif, as well as various PCR-based or low stringency hybridization-based cloning techniques, have advanced the state of the art by making available large numbers of DNA/amino acid sequences for proteins that are known to have biological activity, for example, by virtue of their secreted nature in the case of leader sequence cloning, by virtue of their cell or tissue source in the case of PCR-based techniques, or by virtue of structural similarity to other genes of known biological activity.

Identified polynucleotide and polypeptide sequences have numerous applications in, for example, diagnostics, forensics, gene mapping; identification of mutations responsible for genetic disorders or other traits, to assess biodiversity, and to produce many other types of data and products dependent on DNA and amino acid sequences.

3. SUMMARY OF THE INVENTION

The compositions of the present invention include novel isolated polypeptides, novel isolated polynucleotides encoding such polypeptides, including recombinant DNA molecules, cloned genes or degenerate variants thereof, especially naturally occurring variants such as allelic variants, antisense polynucleotide molecules, and antibodies that specifically recognize one or more epitopes present on such polypeptides, as well as hybridomas producing such antibodies.

The compositions of the present invention additionally include vectors, including expression vectors, containing the polynucleotides of the invention, cells genetically engineered to contain such polynucleotides and cells genetically engineered to express such polynucleotides.

The present invention relates to a collection or library of at least one novel nucleic acid sequence assembled from expressed sequence tags (ESTs) isolated mainly by sequencing by hybridization (SBH), and in some cases, sequences obtained from one or more public databases. The invention relates also to the proteins encoded by such polynucleotides, along with therapeutic, diagnostic and research utilities for these polynucleotides and proteins. These nucleic acid sequences are designated as SEQ ID NO: 1-984, 1969-2952, 3937-3942 or 3949-3954. The polypeptides sequences are designated SEQ ID NO: 985-1968, 2953-3936, 3943-3948 or 3955-3960. The nucleic acids and polypeptides are provided in the Sequence Listing. In the nucleic acids provided in the Sequence Listing, A is adenosine; C is cytosine; G is guanine; T is thymine; and N is any of the four bases. In the amino acids provided in the Sequence Listing, * corresponds to the stop codon.

The nucleic acid sequences of the present invention also include, nucleic acid sequences that hybridize to the complement of SEQ ID NO: 1-984, 1969-2952, 3937-3942 or 3949-3954 under stringent hybridization conditions; nucleic acid sequences which are allelic variants or species homologues of any of the nucleic acid sequences recited above, or nucleic acid sequences that encode a peptide comprising a specific domain or truncation of the peptides encoded by SEQ ID NO: 1-984, 1969-2952, 3937-3942 or 3949-3954. A polynucleotide comprising a nucleotide sequence having at least 90% identity to an identifying sequence of SEQ ID NO: 1-984, 1969-2952, 3937-3942 or 3949-3954 or a degenerate variant or fragment thereof. The identifying sequence can be 100 base pairs in length.

The nucleic acid sequences of the present invention also include the sequence information from the nucleic acid sequences of SEQ ID NO:1-984, 1969-2952, 3937-3942 or 3949-3954. The sequence information can be a segment of any one of SEQ ID NO:1-984, 1969-2952, 3937-3942 or 3949-3954 that uniquely identifies or represents the sequence information of SEQ ID NO:1-984, 1969-2952, 3937-3942 or 3949-3954.

A collection as used in this application can be a collection of only one polynucleotide. The collection of sequence information or identifying information of each sequence can be provided on a nucleic acid array. In one embodiment, segments of sequence information is provided on a nucleic acid array to detect the polynucleotide that contains the segment. The array can be designed to detect full-match or mismatch to the polynucleotide that contains the segment. The collection can also be provided in a computer-readable format.

This invention also includes the reverse or direct complement of any of the nucleic acid sequences recited above; cloning or expression vectors containing the nucleic acid sequences; and host cells or organisms transformed with these expression vectors. Nucleic acid sequences (or their reverse or direct complements) according to the invention have numerous applications in a variety

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of techniques known to those skilled in the art of molecular biology, such as use as hybridization probes, use as primers for PCR, use in an array, use in computer-readable media, use in sequencing full-length genes, use for chromosome and gene mapping, use in the recombinant production of protein, and use in the generation of anti-sense DNA or RNA, their chemical analogs and the like.

In a preferred embodiment, the nucleic acid sequences of SEQ ID NO:1-984, 1969-2952, 3937-3942 or 3949-3954 or novel segments or parts of the nucleic acids of the invention are used as primers in expression assays that are well known in the art. In a particularly preferred embodiment, the nucleic acid sequences of SEQ ID NO:1-984, 1969-2952, 3937-3942 or 3949-3954 or novel segments or parts of the nucleic acids provided herein are used in diagnostics for identifying expressed genes or, as well known in the art and exemplified by Vollrath et al., Science 258:52-59 (1992), as expressed sequence tags for physical mapping of the human genome.

The isolated polynucleotides of the invention include, but are not limited to, a polynucleotide comprising any one of the nucleotide sequences set forth in SEQ ID NO:1-984, 1969-2952, 3937-3942 or 3949-3954; a polynucleotide comprising any of the full length protein coding sequences of SEQ ID NO:1-984, 1969-2952, 3937-3942 or 3949-3954; and a polynucleotide comprising any of the nucleotide sequences of the mature protein coding sequences of SEQ ID NO:1-984, 1969-2952, 3937-3942 or 3949-3954. The polynucleotides of the present invention also include, but are not limited to, a polynucleotide that hybridizes under stringent hybridization conditions to (a) the complement of any one of the nucleotide sequences set forth in SEQ ID NO:1-984, 1969-2952, 3937-3942 or 3949-3954; (b) a nucleotide sequence encoding any one of the amino acid sequences set forth in the Sequence Listing; (c) a polynucleotide which is an allelic variant of any polynucleotides recited above; (d) a polynucleotide which encodes a species homolog (e.g. orthologs) of any of the proteins recited above; or (e) a polynucleotide that encodes a polypeptide comprising a specific domain or truncation of any of the polypeptides comprising an amino acid sequence set forth in the Sequence Listing.

The isolated polypeptides of the invention include, but are not limited to, a polypeptide comprising any of the amino acid sequences set forth in SEQ ID NO: 985-1968, 2953-3936, 3943-3948 or 3955-3960; or the corresponding full length or mature protein. Polypeptides of the invention also include polypeptides with biological activity that are encoded by (a) any of the polynucleotides having a nucleotide sequence set forth in SEQ ID NO:1-984, 1969-2952, 3937-3942 or 3949-3954; or (b) polynucleotides that hybridize to the complement of the polynucleotides of (a) under stringent hybridization conditions. Biologically or immunologically active variants of any of the polypeptide sequences in the Sequence Listing, and "substantial equivalents" thereof (e.g., with at least about 65%, 70%, 75%, 80%, 85%, 90%, 95%, 98% or 99% amino acid sequence identity) that preferably retain biological activity are also contemplated. The polypeptides of the

invention may be wholly or partially chemically synthesized but are preferably produced by recombinant means using the genetically engineered cells (e.g. host cells) of the invention.

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The invention also provides compositions comprising a polypeptide of the invention.

Polypeptide compositions of the invention may further comprise an acceptable carrier, such as a hydrophilic, e.g., pharmaceutically acceptable, carrier.

The invention also provides host cells transformed or transfected with a polynucleotide of the invention.

The invention also relates to methods for producing a polypeptide of the invention comprising growing a culture of the host cells of the invention in a suitable culture medium under conditions permitting expression of the desired polypeptide, and purifying the polypeptide from the culture or from the host cells. Preferred embodiments include those in which the protein produced by such process is a mature form of the protein.

Polynucleotides according to the invention have numerous applications in a variety of techniques known to those skilled in the art of molecular biology. These techniques include use as hybridization probes, use as oligomers, or primers, for PCR, use for chromosome and gene mapping, use in the recombinant production of protein, and use in generation of anti-sense DNA or RNA, their chemical analogs and the like. For example, when the expression of an mRNA is largely restricted to a particular cell or tissue type, polynucleotides of the invention can be used as hybridization probes to detect the presence of the particular cell or tissue mRNA in a sample using, e.g., in situ hybridization.

In other exemplary embodiments, the polynucleotides are used in diagnostics as expressed sequence tags for identifying expressed genes or, as well known in the art and exemplified by Vollrath et al., Science 258:52-59 (1992), as expressed sequence tags for physical mapping of the human genome.

The polypeptides according to the invention can be used in a variety of conventional procedures and methods that are currently applied to other proteins. For example, a polypeptide of the invention can be used to generate an antibody that specifically binds the polypeptide. Such antibodies, particularly monoclonal antibodies, are useful for detecting or quantitating the polypeptide in tissue. The polypeptides of the invention can also be used as molecular weight markers, and as a food supplement.

Methods are also provided for preventing, treating, or ameliorating a medical condition which comprises the step of administering to a mammalian subject a therapeutically effective amount of a composition comprising a polypeptide of the present invention and a pharmaceutically acceptable carrier.

In particular, the polypeptides and polynucleotides of the invention can be utilized, for example, in methods for the prevention and/or treatment of disorders involving aberrant protein expression or biological activity.

The present invention further relates to methods for detecting the presence of the polynucleotides or polypeptides of the invention in a sample. Such methods can, for example, be utilized as part of prognostic and diagnostic evaluation of disorders as recited herein and for the identification of subjects exhibiting a predisposition to such conditions. The invention provides a method for detecting the polynucleotides of the invention in a sample, comprising contacting the sample with a compound that binds to and forms a complex with the polynucleotide of interest for a period sufficient to form the complex and under conditions sufficient to form a complex and detecting the complex such that if a complex is detected, the polynucleotide of interest is detected. The invention also provides a method for detecting the polypeptides of the invention in a sample comprising contacting the sample with a compound that binds to and forms a complex with the polypeptide under conditions and for a period sufficient to form the complex and detecting the formation of the complex such that if a complex is formed, the polypeptide is detected.

The invention also provides kits comprising polynucleotide probes and/or monoclonal antibodies, and optionally quantitative standards, for carrying out methods of the invention. Furthermore, the invention provides methods for evaluating the efficacy of drugs, and monitoring the progress of patients, involved in clinical trials for the treatment of disorders as recited above.

The invention also provides methods for the identification of compounds that modulate (i.e., increase or decrease) the expression or activity of the polynucleotides and/or polypeptides of the invention. Such methods can be utilized, for example, for the identification of compounds that can ameliorate symptoms of disorders as recited herein. Such methods can include, but are not limited to, assays for identifying compounds and other substances that interact with (e.g., bind to) the polypeptides of the invention. The invention provides a method for identifying a compound that binds to the polypeptides of the invention comprising contacting the compound with a polypeptide of the invention in a cell for a time sufficient to form a polypeptide/compound complex, wherein the complex drives expression of a reporter gene sequence in the cell; and detecting the complex by detecting the reporter gene sequence expression such that if expression of the reporter gene is detected the compound the binds to a polypeptide of the invention is identified.

The methods of the invention also provides methods for treatment which involve the administration of the polynucleotides or polypeptides of the invention to individuals exhibiting

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symptoms or tendencies. In addition, the invention encompasses methods for treating diseases or disorders as recited herein comprising administering compounds and other substances that modulate the overall activity of the target gene products. Compounds and other substances can effect such modulation either on the level of target gene/protein expression or target protein activity.

The polypeptides of the present invention and the polynucleotides encoding them are also useful for the same functions known to one of skill in the art as the polypeptides and polynucleotides to which they have homology (set forth in Tables 2 and 9); for which they have a signature region (as set forth in Tables 3 and 10); or for which they have homology to a gene family (as set forth in Tables 4 and 11). If no homology is set forth for a sequence, then the polypeptides and polynucleotides of the present invention are useful for a variety of applications, as described herein, including use in arrays for detection.

4. DETAILED DESCRIPTION OF THE INVENTION

4.1 DEFINITIONS

It must be noted that as used herein and in the appended claims, the singular forms "a", "an" and "the" include plural references unless the context clearly dictates otherwise.

The term "active" refers to those forms of the polypeptide which retain the biologic and/or immunologic activities of any naturally occurring polypeptide. According to the invention, the terms "biologically active" or "biological activity" refer to a protein or peptide having structural, regulatory or biochemical functions of a naturally occurring molecule. Likewise "immunologically active" or "immunological activity" refers to the capability of the natural, recombinant or synthetic polypeptide to induce a specific immune response in appropriate animals or cells and to bind with specific antibodies.

The term "activated cells" as used in this application are those cells which are engaged in extracellular or intracellular membrane trafficking, including the export of secretory or enzymatic molecules as part of a normal or disease process.

The terms "complementary" or "complementarity" refer to the natural binding of polynucleotides by base pairing. For example, the sequence 5'-AGT-3' binds to the complementary sequence 3'-TCA-5'. Complementarity between two single-stranded molecules may be "partial" such that only some of the nucleic acids bind or it may be "complete" such that total complementarity exists between the single stranded molecules. The degree of complementarity between the nucleic acid strands has significant effects on the efficiency and strength of the hybridization between the nucleic acid strands.

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The term "embryonic stem cells (ES)" refers to a cell that can give rise to many differentiated cell types in an embryo or an adult, including the germ cells. The term "germ line stem cells (GSCs)" refers to stem cells derived from primordial stem cells that provide a steady and continuous source of germ cells for the production of gametes. The term "primordial germ

cells (PGCs)" refers to a small population of cells set aside from other cell lineages particularly from the yolk sac, mesenteries, or gonadal ridges during embryogenesis that have the potential to differentiate into germ cells and other cells. PGCs are the source from which GSCs and ES cells are derived The PGCs, the GSCs and the ES cells are capable of self-renewal. Thus these cells not only populate the germ line and give rise to a plurality of terminally differentiated cells that comprise the adult specialized organs, but are able to regenerate themselves.

The term "expression modulating fragment," EMF, means a series of nucleotides which modulates the expression of an operably linked ORF or another EMF.

As used herein, a sequence is said to "modulate the expression of an operably linked sequence" when the expression of the sequence is altered by the presence of the EMF. EMFs include, but are not limited to, promoters, and promoter modulating sequences (inducible elements). One class of EMFs are nucleic acid fragments which induce the expression of an operably linked ORF in response to a specific regulatory factor or physiological event.

The terms "nucleotide sequence" or "nucleic acid" or "polynucleotide" or "oligonculeotide" are used interchangeably and refer to a heteropolymer of nucleotides or the sequence of these nucleotides. These phrases also refer to DNA or RNA of genomic or synthetic origin which may be single-stranded or double-stranded and may represent the sense or the antisense strand, to peptide nucleic acid (PNA) or to any DNA-like or RNA-like material. In the sequences herein A is adenine, C is cytosine, T is thymine, G is guanine and N is A, C, G or T (U). It is contemplated that where the polynucleotide is RNA, the T (thymine) in the sequences provided herein is substituted with U (uracil). Generally, nucleic acid segments provided by this invention may be assembled from fragments of the genome and short oligonucleotide linkers, or from a series of oligonucleotides, or from individual nucleotides, to provide a synthetic nucleic acid which is capable of being expressed in a recombinant transcriptional unit comprising regulatory elements derived from a microbial or viral operon, or a eukaryotic gene.

The terms "oligonucleotide fragment" or a "polynucleotide fragment", "portion," or "segment" or "probe" or "primer" are used interchangeably and refer to a sequence of nucleotide residues which are at least about 5 nucleotides, more preferably at least about 7 nucleotides, more preferably at least about 11 nucleotides and most preferably at least about 17 nucleotides. The fragment is preferably less than about 500 nucleotides, preferably less than about 200 nucleotides, more preferably less than about 100

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nucleotides, more preferably less than about 50 nucleotides and most preferably less than 30 nucleotides. Preferably the probe is from about 6 nucleotides to about 200 nucleotides, preferably from about 15 to about 50 nucleotides, more preferably from about 17 to 30 nucleotides and most preferably from about 20 to 25 nucleotides. Preferably the fragments can be used in polymerase chain reaction (PCR), various hybridization procedures or microarray procedures to identify or amplify identical or related parts of mRNA or DNA molecules. A fragment or segment may uniquely identify each polynucleotide sequence of the present invention. Preferably the fragment comprises a sequence substantially similar to any one of SEQ ID NOs:1-20.

Probes may, for example, be used to determine whether specific mRNA molecules are present in a cell or tissue or to isolate similar nucleic acid sequences from chromosomal DNA as described by Walsh et al. (Walsh, P.S. et al., 1992, PCR Methods Appl 1:241-250). They may be labeled by nick translation, Klenow fill-in reaction, PCR, or other methods well known in the art. Probes of the present invention, their preparation and/or labeling are elaborated in Sambrook, J. et al., 1989, Molecular Cloning: A Laboratory Manual, Cold Spring Harbor Laboratory, NY; or Ausubel, F.M. et al., 1989, Current Protocols in Molecular Biology, John Wiley & Sons, New York NY, both of which are incorporated herein by reference in their entirety.

The nucleic acid sequences of the present invention also include the sequence information from the nucleic acid sequences of SEQ ID NO: 1-984, 1969-2952, 3937-3942 or 3949-3954. The sequence information can be a segment of any one of SEQ ID NO:1-1-984, 1969-2952, 3937-3942 or 3949-3954 that uniquely identifies or represents the sequence information of that sequence of SEQ ID NO:1-984, 1969-2952, 3937-3942 or 3949-3954. One such segment can be a twenty-mer nucleic acid sequence because the probability that a twenty-mer is fully matched in the human genome is 1 in 300. In the human genome, there are three billion base pairs in one set of chromosomes. Because 4²⁰ possible twenty-mers exist, there are 300 times more twenty-mers than there are base pairs in a set of human chromosomes. Using the same analysis, the probability for a seventeen-mer to be fully matched in the human genome is approximately 1 in 5. When these segments are used in arrays for expression studies, fifteenmer segments can be used. The probability that the fifteen-mer is fully matched in the expressed sequences is also approximately one in five because expressed sequences comprise less than approximately 5% of the entire genome sequence.

Similarly, when using sequence information for detecting a single mismatch, a segment can be a twenty-five mer. The probability that the twenty-five mer would appear in a human genome with a single mismatch is calculated by multiplying the probability for a full match (1÷4²⁵) times the

increased probability for mismatch at each nucleotide position (3 x 25). The probability that an eighteen mer with a single mismatch can be detected in an array for expression studies is approximately one in five. The probability that a twenty-mer with a single mismatch can be

detected in a human genome is approximately one in five.

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The term "open reading frame," ORF, means a series of nucleotide triplets coding for amino acids without any termination codons and is a sequence translatable into protein.

The terms "operably linked" or "operably associated" refer to functionally related nucleic acid sequences. For example, a promoter is operably associated or operably linked with a coding sequence if the promoter controls the transcription of the coding sequence. While operably linked nucleic acid sequences can be contiguous and in the same reading frame, certain genetic elements e.g. repressor genes are not contiguously linked to the coding sequence but still control transcription/translation of the coding sequence.

The term "pluripotent" refers to the capability of a cell to differentiate into a number of differentiated cell types that are present in an adult organism. A pluripotent cell is restricted in its differentiation capability in comparison to a totipotent cell.

The terms "polypeptide" or "peptide" or "amino acid sequence" refer to an oligopeptide, peptide, polypeptide or protein sequence or fragment thereof and to naturally occurring or synthetic molecules. A polypeptide "fragment," "portion," or "segment" is a stretch of amino acid residues of at least about 5 amino acids, preferably at least about 7 amino acids, more preferably at least about 9 amino acids and most preferably at least about 17 or more amino acids. The peptide preferably is not greater than about 500 amino acids, more preferably less than 200 amino acids more preferably less than 150 amino acids and most preferably less than 100 amino acids. Preferably the peptide is from about 5 to about 200 amino acids. To be active, any polypeptide must have sufficient length to display biological and/or immunological activity.

The term "naturally occurring polypeptide" refers to polypeptides produced by cells that have not been genetically engineered and specifically contemplates various polypeptides arising from post-translational modifications of the polypeptide including, but not limited to, acetylation, carboxylation, glycosylation, phosphorylation, lipidation and acylation.

The term "translated protein coding portion" means a sequence which encodes for the full length protein which may include any leader sequence or any processing sequence.

The term "mature protein coding sequence" means a sequence which encodes a peptide or protein without a signal or leader sequence. The "mature protein portion" means that portion of the protein which does not include a signal or leader sequence. The peptide may have been produced by processing in the cell which removes any leader/signal sequence. The mature protein portion may or may not include the initial methionine residue. The methionine residue

may be removed from the protein during processing in the cell. The peptide may be produced synthetically or the protein may have been produced using a polynucleotide only encoding for the mature protein coding sequence.

The term "derivative" refers to polypeptides chemically modified by such techniques as ubiquitination, labeling (e.g., with radionuclides or various enzymes), covalent polymer attachment such as pegylation (derivatization with polyethylene glycol) and insertion or substitution by chemical synthesis of amino acids such as ornithine, which do not normally occur in human proteins.

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The term "variant" (or "analog") refers to any polypeptide differing from naturally occurring polypeptides by amino acid insertions, deletions, and substitutions, created using, e.g., recombinant DNA techniques. Guidance in determining which amino acid residues may be replaced, added or deleted without abolishing activities of interest, may be found by comparing the sequence of the particular polypeptide with that of homologous peptides and minimizing the number of amino acid sequence changes made in regions of high homology (conserved regions) or by replacing amino acids with consensus sequence.

Alternatively, recombinant variants encoding these same or similar polypeptides may be synthesized or selected by making use of the "redundancy" in the genetic code. Various codon substitutions, such as the silent changes which produce various restriction sites, may be introduced to optimize cloning into a plasmid or viral vector or expression in a particular prokaryotic or eukaryotic system. Mutations in the polypucleotide sequence may be reflected in the polypeptide or domains of other peptides added to the polypeptide to modify the properties of any part of the polypeptide, to change characteristics such as ligand-binding affinities, interchain affinities, or degradation/turnover rate.

Preferably, amino acid "substitutions" are the result of replacing one amino acid with another amino acid having similar structural and/or chemical properties, *i.e.*, conservative amino acid replacements. "Conservative" amino acid substitutions may be made on the basis of similarity in polarity, charge, solubility, hydrophobicity, hydrophilicity, and/or the amphipathic nature of the residues involved. For example, nonpolar (hydrophobic) amino acids include alanine, leucine, isoleucine, valine, proline, phenylalanine, tryptophan, and methionine; polar neutral amino acids include glycine, serine, threonine, cysteine, tyrosine, asparagine, and glutamine; positively charged (basic) amino acids include arginine, lysine, and histidine; and negatively charged (acidic) amino acids include aspartic acid and glutamic acid. "Insertions" or "deletions" are preferably in the range of about 1 to 20 amino acids, more preferably 1 to 10 amino acids. The variation allowed may be experimentally determined by systematically making

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insertions, deletions, or substitutions of amino acids in a polypeptide molecule using recombinant DNA techniques and assaying the resulting recombinant variants for activity.

Alternatively, where alteration of function is desired, insertions, deletions or non-conservative alterations can be engineered to produce altered polypeptides. Such alterations can, for example, alter one or more of the biological functions or biochemical characteristics of the polypeptides of the invention. For example, such alterations may change polypeptide characteristics such as ligand-binding affinities, interchain affinities, or degradation/turnover rate. Further, such alterations can be selected so as to generate polypeptides that are better suited for expression, scale up and the like in the host cells chosen for expression. For example, cysteine residues can be deleted or substituted with another amino acid residue in order to eliminate disulfide bridges.

The terms "purified" or "substantially purified" as used herein denotes that the indicated nucleic acid or polypeptide is present in the substantial absence of other biological macromolecules, e.g., polynucleotides, proteins, and the like. In one embodiment, the polynucleotide or polypeptide is purified such that it constitutes at least 95% by weight, more preferably at least 99% by weight, of the indicated biological macromolecules present (but water, buffers, and other small molecules, especially molecules having a molecular weight of less than 1000 daltons, can be present).

The term "isolated" as used herein refers to a nucleic acid or polypeptide separated from at least one other component (e.g., nucleic acid or polypeptide) present with the nucleic acid or polypeptide in its natural source. In one embodiment, the nucleic acid or polypeptide is found in the presence of (if anything) only a solvent, buffer, ion, or other component normally present in a solution of the same. The terms "isolated" and "purified" do not encompass nucleic acids or polypeptides present in their natural source.

The term "recombinant," when used herein to refer to a polypeptide or protein, means that a polypeptide or protein is derived from recombinant (e.g., microbial, insect, or mammalian) expression systems. "Microbial" refers to recombinant polypeptides or proteins made in bacterial or fungal (e.g., yeast) expression systems. As a product, "recombinant microbial" defines a polypeptide or protein essentially free of native endogenous substances and unaccompanied by associated native glycosylation. Polypeptides or proteins expressed in most bacterial cultures, e.g., E. coli, will be free of glycosylation modifications; polypeptides or proteins expressed in yeast will have a glycosylation pattern in general different from those expressed in mammalian cells.

The term "recombinant expression vehicle or vector" refers to a plasmid or phage or virus or vector, for expressing a polypeptide from a DNA (RNA) sequence. An expression vehicle can

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comprise a transcriptional unit comprising an assembly of (1) a genetic element or elements having a regulatory role in gene expression, for example, promoters or enhancers, (2) a structural or coding sequence which is transcribed into mRNA and translated into protein, and (3) appropriate transcription initiation and termination sequences. Structural units intended for use in yeast or eukaryotic expression systems preferably include a leader sequence enabling extracellular secretion of translated protein by a host cell. Alternatively, where recombinant protein is expressed without a leader or transport sequence, it may include an amino terminal methionine residue. This residue may or may not be subsequently cleaved from the expressed recombinant protein to provide a final product.

The term "recombinant expression system" means host cells which have stably integrated a recombinant transcriptional unit into chromosomal DNA or carry the recombinant transcriptional unit extrachromosomally. Recombinant expression systems as defined herein will express heterologous polypeptides or proteins upon induction of the regulatory elements linked to the DNA segment or synthetic gene to be expressed. This term also means host cells which have stably integrated a recombinant genetic element or elements having a regulatory role in gene expression, for example, promoters or enhancers. Recombinant expression systems as defined herein will express polypeptides or proteins endogenous to the cell upon induction of the regulatory elements linked to the endogenous DNA segment or gene to be expressed. The cells can be prokaryotic or eukaryotic.

The term "secreted" includes a protein that is transported across or through a membrane, including transport as a result of signal sequences in its amino acid sequence when it is expressed in a suitable host cell. "Secreted" proteins include without limitation proteins secreted wholly (e.g., soluble proteins) or partially (e.g., receptors) from the cell in which they are expressed. "Secreted" proteins also include without limitation proteins that are transported across the membrane of the endoplasmic reticulum. "Secreted" proteins are also intended to include proteins containing non-typical signal sequences (e.g. Interleukin-1 Beta, see Krasney, P.A. and Young, P.R. (1992) Cytokine 4(2):134-143) and factors released from damaged cells (e.g. Interleukin-1 Receptor Antagonist, see Arend, W.P. et. al. (1998) Annu. Rev. Immunol. 16:27-55)

Where desired, an expression vector may be designed to contain a "signal or leader sequence" which will direct the polypeptide through the membrane of a cell. Such a sequence may be naturally present on the polypeptides of the present invention or provided from heterologous protein sources by recombinant DNA techniques.

The term "stringent" is used to refer to conditions that are commonly understood in the art as stringent. Stringent conditions can include highly stringent conditions (i.e., hybridization

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to filter-bound DNA in 0.5 M NaHPO₄, 7% sodium dodecyl sulfate (SDS), 1 mM EDTA at 65°C, and washing in 0.1X SSC/0.1% SDS at 68°C), and moderately stringent conditions (i.e., washing in 0.2X SSC/0.1% SDS at 42°C). Other exemplary hybridization conditions are described herein in the examples.

In instances of hybridization of deoxyoligonucleotides, additional exemplary stringent hybridization conditions include washing in 6X SSC/0.05% sodium pyrophosphate at 37°C (for 14-base oligonucleotides), 48°C (for 17-base oligos), 55°C (for 20-base oligonucleotides), and 60°C (for 23-base oligonucleotides).

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As used herein, "substantially equivalent" can refer both to nucleotide and amino acid sequences, for example a mutant sequence, that varies from a reference sequence by one or more substitutions, deletions, or additions, the net effect of which does not result in an adverse functional dissimilarity between the reference and subject sequences. Typically, such a substantially equivalent sequence varies from one of those listed herein by no more than about 35% (i.e., the number of individual residue substitutions, additions, and/or deletions in a substantially equivalent sequence, as compared to the corresponding reference sequence, divided by the total number of residues in the substantially equivalent sequence is about 0.35 or less). Such a sequence is said to have 65% sequence identity to the listed sequence. In one embodiment, a substantially equivalent, e.g., mutant, sequence of the invention varies from a listed sequence by no more than 30% (70% sequence identity); in a variation of this embodiment. by no more than 25% (75% sequence identity); and in a further variation of this embodiment, by no more than 20% (80% sequence identity) and in a further variation of this embodiment, by no more than 10% (90% sequence identity) and in a further variation of this embodiment, by no more that 5% (95% sequence identity). Substantially equivalent, e.g., mutant, amino acid sequences according to the invention preferably have at least 80% sequence identity with a listed amino acid sequence, more preferably at least 85% sequence identity, more preferably at least 90% sequence identity, more preferably at least 95% sequence identity, more preferably at least 98% sequence identity and most preferably at least 98% idenity. Substantially equivalent nucleotide sequences of the invention can have lower percent sequence identities, taking into account, for example, the redundancy or degeneracy of the genetic code. Preferably, nucleotide sequence has at least about 65% identity, more preferably at least about 75% identity, more preferably at least about 80% identity, more preferably at least about 85% identity, more preferably at least about 90% identity, and most preferably at least about 95% identity, more preferably at least 98% and most preferably at least about 99% identity. For the purposes of the present invention, sequences having substantially equivalent biological activity and substantially equivalent expression characteristics are considered substantially equivalent. For the purposes of

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determining equivalence, truncation of the mature sequence (e.g., via a mutation which creates a spurious stop codon) should be disregarded. Sequence identity may be determined, e.g., using the Jotun Hein method (Hein, J. (1990) Methods Enzymol. 183:626-645). Identity between sequences can also be determined by other methods known in the art, e.g. by varying hybridization conditions.

The term "totipotent" refers to the capability of a cell to differentiate into all of the cell types of an adult organism.

The term "transformation" means introducing DNA into a suitable host cell so that the DNA is replicable, either as an extrachromosomal element, or by chromosomal integration. The term "transfection" refers to the taking up of an expression vector by a suitable host cell, whether or not any coding sequences are in fact expressed. The term "infection" refers to the introduction of nucleic acids into a suitable host cell by use of a virus or viral vector.

As used herein, an "uptake modulating fragment," UMF, means a series of nucleotides which mediate the uptake of a linked DNA fragment into a cell. UMFs can be readily identified using known UMFs as a target sequence or target motif with the computer-based systems described below. The presence and activity of a UMF can be confirmed by attaching the suspected UMF to a marker sequence. The resulting nucleic acid molecule is then incubated with an appropriate host under appropriate conditions and the uptake of the marker sequence is determined. As described above, a UMF will increase the frequency of uptake of a linked marker sequence.

Each of the above terms is meant to encompass all that is described for each, unless the context dictates otherwise.

4.2 NUCLEIC ACIDS OF THE INVENTION

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Nucleotide sequences of the invention are set forth in the Sequence Listing.

The isolated polynucleotides of the invention include a polynucleotide comprising the nucleotide sequences of SEQ ID NO: 1-984, 1969-2952, 3937-3942 or 3949-3954; a polynucleotide encoding any one of the peptide sequences of SEQ ID NO: 985-1968, 2953-3936, 3943-3948 or 3955-3960; and a polynucleotide comprising the nucleotide sequence encoding the mature protein coding sequence of the polypeptides of any one of SEQ ID NO: 985-1968, 2953-3936, 3943-3948 or 3955-3960. The polynucleotides of the present invention also include, but are not limited to, a polynucleotide that hybridizes under stringent conditions to (a) the complement of any of the nucleotides sequences of SEQ ID NO: 1-984, 1969-2952, 3937-3942 or 3949-3954; (b) nucleotide sequences encoding any one of the amino acid sequences set forth in the Sequence Listing as SEQ ID NO: 985-1968, 2953-3936, 3943-3948 or 3955-3960; (c) a

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polynucleotide which is an allelic variant of any polynucleotide recited above; (d) a polynucleotide which encodes a species homolog of any of the proteins recited above; or (e) a polynucleotide that encodes a polypeptide comprising a specific domain or truncation of the polypeptides of SEQ ID NO:985-1968, 2953-3936, 3943-3948 or 3955-3960. Domains of interest may depend on the nature of the encoded polypeptide; e.g., domains in receptor-like polypeptides include ligand-binding, extracellular, transmembrane, or cytoplasmic domains, or combinations thereof; domains in immunoglobulin-like proteins include the variable immunoglobulin-like domains; domains in enzyme-like polypeptides include catalytic and substrate binding domains; and domains in ligand polypeptides include receptor-binding domains.

The polynucleotides of the invention include naturally occurring or wholly or partially synthetic DNA, e.g., cDNA and genomic DNA, and RNA, e.g., mRNA. The polynucleotides may include all of the coding region of the cDNA or may represent a portion of the coding region of the cDNA.

The present invention also provides genes corresponding to the cDNA sequences disclosed herein. The corresponding genes can be isolated in accordance with known methods using the sequence information disclosed herein. Such methods include the preparation of probes or primers from the disclosed sequence information for identification and/or amplification of genes in appropriate genomic libraries or other sources of genomic materials. Further 5' and 3' sequence can be obtained using methods known in the art. For example, full length cDNA or genomic DNA that corresponds to any of the polynucleotides of SEQ ID NO: 1-984, 1969-2952, 3937-3942 or 3949-3954 can be obtained by screening appropriate cDNA or genomic DNA libraries under suitable hybridization conditions using any of the polynucleotides of SEQ ID NO: 1-984, 1969-2952, 3937-3942 or 3949-3954 or a portion thereof as a probe. Alternatively, the polynucleotides of SEQ ID NO: 1-984, 1969-2952, 3937-3942 or 3949-3954 may be used as the basis for suitable primer(s) that allow identification and/or amplification of genes in appropriate genomic DNA or cDNA libraries.

The nucleic acid sequences of the invention can be assembled from ESTs and sequences (including cDNA and genomic sequences) obtained from one or more public databases, such as dbEST, gbpri, and UniGene. The EST sequences can provide identifying sequence information, representative fragment or segment information, or novel segment information for the full-length gene.

The polynucleotides of the invention also provide polynucleotides including nucleotide sequences that are substantially equivalent to the polynucleotides recited above. Polynucleotides according to the invention can have, e.g., at least about 65%, at least about 70%, at least about

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75%, at least about 80%, 81%, 82%, 83%, 84%, more typically at least about 85%, 86%, 87%, 88%, 89%, and more typically at least about 90%, 91%, 92%, 93%, 94%, and even more typically at least about 95%, 96%, 97%, 98%, 99%, sequence identity to a polynucleotide recited above.

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Included within the scope of the nucleic acid sequences of the invention are nucleic acid sequence fragments that hybridize under stringent conditions to any of the nucleotide sequences of SEQ ID NO: 1-984, 1969-2952, 3937-3942 or 3949-3954, or complements thereof, which fragment is greater than about 5 nucleotides, preferably 7 nucleotides, more preferably greater than 9 nucleotides and most preferably greater than 17 nucleotides. Fragments of, e.g. 15, 17, or 20 nucleotides or more that are selective for (i.e. specifically hybridize to any one of the polynucleotides of the invention) are contemplated. Probes capable of specifically hybridizing to a polynucleotide can differentiate polynucleotide sequences of the invention from other polynucleotide sequences in the same family of genes or can differentiate human genes from genes of other species, and are preferably based on unique nucleotide sequences.

The sequences falling within the scope of the present invention are not limited to these specific sequences, but also include allelic and species variations thereof. Allelic and species variations can be routinely determined by comparing the sequence provided SEQ ID NO: 1-984, 1969-2952, 3937-3942 or 3949-3954, a representative fragment thereof, or a nucleotide sequence at least 90% identical, preferably 95% identical, to SEQ ID NO: 1-984, 1969-2952, 3937-3942 or 3949-3954 with a sequence from another isolate of the same species. Furthermore, to accommodate codon variability, the invention includes nucleic acid molecules coding for the same amino acid sequences as do the specific ORFs disclosed herein. In other words, in the coding region of an ORF, substitution of one codon for another codon that encodes the same amino acid is expressly contemplated.

The nearest neighbor or homology result for the nucleic acids of the present invention, including SEQ ID NO: 1-984, 1969-2952, 3937-3942 or 3949-3954, can be obtained by searching a database using an algorithm or a program. Preferably, a BLAST which stands for Basic Local Alignment Search Tool is used to search for local sequence alignments (Altshul, S.F. J Mol. Evol. 36 290-300 (1993) and Altschul S.F. et al. J. Mol. Biol. 21:403-410 (1990)). Alternatively a FASTA version 3 search against Genpept, using Fastxy algorithm.

Species homologs (or orthologs) of the disclosed polynucleotides and proteins are also provided by the present invention. Species homologs may be isolated and identified by making suitable probes or primers from the sequences provided herein and screening a suitable nucleic acid source from the desired species.

The invention also encompasses allelic variants of the disclosed polynucleotides or proteins; that is, naturally-occurring alternative forms of the isolated polynucleotide which also encode proteins which are identical, homologous or related to that encoded by the polynucleotides.

The nucleic acid sequences of the invention are further directed to sequences which encode variants of the described nucleic acids. These amino acid sequence variants may be prepared by methods known in the art by introducing appropriate nucleotide changes into a native or variant polynucleotide. There are two variables in the construction of amino acid sequence variants: the location of the mutation and the nature of the mutation. Nucleic acids encoding the amino acid sequence variants are preferably constructed by mutating the polynucleotide to encode an amino acid sequence that does not occur in nature. These nucleic acid alterations can be made at sites that differ in the nucleic acids from different species (variable positions) or in highly conserved regions (constant regions). Sites at such locations will typically be modified in series, e.g., by substituting first with conservative choices (e.g., hydrophobic amino acid to a different hydrophobic amino acid) and then with more distant choices (e.g., hydrophobic amino acid to a charged amino acid), and then deletions or insertions may be made at the target site. Amino acid sequence deletions generally range from about 1 to 30 residues, preferably about 1 to 10 residues, and are typically contiguous. Amino acid insertions include amino- and/or carboxyl-terminal fusions ranging in length from one to one hundred or more residues, as well as intrasequence insertions of single or multiple amino acid residues. Intrasequence insertions may range generally from about 1 to 10 amino residues, preferably from 1 to 5 residues. Examples of terminal insertions include the heterologous signal sequences necessary for secretion or for intracellular targeting in different host cells and sequences such as FLAG or poly-histidine sequences useful for purifying the expressed protein.

In a preferred method, polynucleotides encoding the novel amino acid sequences are changed via site-directed mutagenesis. This method uses oligonucleotide sequences to alter a polynucleotide to encode the desired amino acid variant, as well as sufficient adjacent nucleotides on both sides of the changed amino acid to form a stable duplex on either side of the site of being changed. In general, the techniques of site-directed mutagenesis are well known to those of skill in the art and this technique is exemplified by publications such as, Edelman et al., DNA 2:183 (1983). A versatile and efficient method for producing site-specific changes in a polynucleotide sequence was published by Zoller and Smith, Nucleic Acids Res. 10:6487-6500 (1982). PCR may also be used to create amino acid sequence variants of the novel nucleic acids. When small amounts of template DNA are used as starting material, primer(s) that differs slightly in sequence from the corresponding region in the template DNA can generate the desired

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amino acid variant. PCR amplification results in a population of product DNA fragments that differ from the polynucleotide template encoding the polypeptide at the position specified by the primer. The product DNA fragments replace the corresponding region in the plasmid and this gives a polynucleotide encoding the desired amino acid variant.

A further technique for generating amino acid variants is the cassette mutagenesis technique described in Wells et al., *Gene* 34:315 (1985); and other mutagenesis techniques well known in the art, such as, for example, the techniques in Sambrook et al., supra, and *Current Protocols in Molecular Biology*, Ausubel et al. Due to the inherent degeneracy of the genetic code, other DNA sequences which encode substantially the same or a functionally equivalent amino acid sequence may be used in the practice of the invention for the cloning and expression of these novel nucleic acids. Such DNA sequences include those which are capable of hybridizing to the appropriate novel nucleic acid sequence under stringent conditions.

Polynucleotides encoding preferred polypeptide truncations of the invention can be used to generate polynucleotides encoding chimeric or fusion proteins comprising one or more domains of the invention and heterologous protein sequences.

The polynucleotides of the invention additionally include the complement of any of the polynucleotides recited above. The polynucleotide can be DNA (genomic, cDNA, amplified, or synthetic) or RNA. Methods and algorithms for obtaining such polynucleotides are well known to those of skill in the art and can include, for example, methods for determining hybridization conditions that can routinely isolate polynucleotides of the desired sequence identities.

In accordance with the invention, polynucleotide sequences comprising the mature protein coding sequences corresponding to any one of SEQ ID NO: 1-984, 1969-2952, 3937-3942 or 3949-3954, or functional equivalents thereof, may be used to generate recombinant DNA molecules that direct the expression of that nucleic acid, or a functional equivalent thereof, in appropriate host cells. Also included are the cDNA inserts of any of the clones identified herein.

A polynucleotide according to the invention can be joined to any of a variety of other nucleotide sequences by well-established recombinant DNA techniques (see Sambrook J et al. (1989) Molecular Cloning: A Laboratory Manual, Cold Spring Harbor Laboratory, NY). Useful nucleotide sequences for joining to polynucleotides include an assortment of vectors, e.g., plasmids, cosmids, lambda phage derivatives, phagemids, and the like, that are well known in the art. Accordingly, the invention also provides a vector including a polynucleotide of the invention and a host cell containing the polynucleotide. In general, the vector contains an origin of replication functional in at least one organism, convenient restriction endonuclease sites, and a selectable marker for the host cell. Vectors according to the invention include expression

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vectors, replication vectors, probe generation vectors, and sequencing vectors. A host cell according to the invention can be a prokaryotic or eukaryotic cell and can be a unicellular organism or part of a multicellular organism.

The present invention further provides recombinant constructs comprising a nucleic acid having any of the nucleotide sequences of SEQ ID NO: 1-984, 1969-2952, 3937-3942 or 3949-3954 or a fragment thereof or any other polynucleotides of the invention. In one embodiment, the recombinant constructs of the present invention comprise a vector, such as a plasmid or viral vector, into which a nucleic acid having any of the nucleotide sequences of SEQ ID NO: 1-984, 1969-2952, 3937-3942 or 3949-3954 or a fragment thereof is inserted, in a forward or reverse orientation. In the case of a vector comprising one of the ORFs of the present invention, the vector may further comprise regulatory sequences, including for example, a promoter, operably linked to the ORF. Large numbers of suitable vectors and promoters are known to those of skill in the art and are commercially available for generating the recombinant constructs of the present invention. The following vectors are provided by way of example. Bacterial: pBs, phagescript, PsiX174, pBluescript SK, pBs KS, pNH8a, pNH16a, pNH18a, pNH46a (Stratagene); pTrc99A, pKK223-3, pKK233-3, pDR540, pRIT5 (Pharmacia). Eukaryotic: pWLneo, pSV2cat, pOG44, PXTI, pSG (Stratagene) pSVK3, pBPV, pMSG, pSVL (Pharmacia).

The isolated polynucleotide of the invention may be operably linked to an expression control sequence such as the pMT2 or pED expression vectors disclosed in Kaufman et al., *Nucleic Acids Res.* 19, 4485-4490 (1991), in order to produce the protein recombinantly. Many suitable expression control sequences are known in the art. General methods of expressing recombinant proteins are also known and are exemplified in R. Kaufman, *Methods in Enzymology* 185, 537-566 (1990). As defined herein "operably linked" means that the isolated polynucleotide of the invention and an expression control sequence are situated within a vector or cell in such a way that the protein is expressed by a host cell which has been transformed (transfected) with the ligated polynucleotide/expression control sequence.

Promoter regions can be selected from any desired gene using CAT (chloramphenicol transferase) vectors or other vectors with selectable markers. Two appropriate vectors are pKK232-8 and pCM7. Particular named bacterial promoters include lacI, lacZ, T3, T7, gpt, lambda PR, and trc. Eukaryotic promoters include CMV immediate early, HSV thymidine kinase, early and late SV40, LTRs from retrovirus, and mouse metallothionein-I. Selection of the appropriate vector and promoter is well within the level of ordinary skill in the art. Generally, recombinant expression vectors will include origins of replication and selectable markers permitting transformation of the host cell, e.g., the ampicillin resistance gene of E. coliand S. cerevisiae TRP1 gene, and a promoter derived from a highly-expressed gene to direct

transcription of a downstream structural sequence. Such promoters can be derived from operons encoding glycolytic enzymes such as 3-phosphoglycerate kinase (PGK), a-factor, acid phosphatase, or heat shock proteins, among others. The heterologous structural sequence is assembled in appropriate phase with translation initiation and termination sequences, and preferably, a leader sequence capable of directing secretion of translated protein into the periplasmic space or extracellular medium. Optionally, the heterologous sequence can encode a fusion protein including an amino terminal identification peptide imparting desired characteristics, e.g., stabilization or simplified purification of expressed recombinant product. Useful expression vectors for bacterial use are constructed by inserting a structural DNA sequence encoding a desired protein together with suitable translation initiation and termination signals in operable reading phase with a functional promoter. The vector will comprise one or more phenotypic selectable markers and an origin of replication to ensure maintenance of the vector and to, if desirable, provide amplification within the host. Suitable prokaryotic hosts for transformation include E. coli, Bacillus subtilis, Salmonella typhimurium and various species within the genera Pseudomonas, Streptomyces, and Staphylococcus, although others may also be employed as a matter of choice.

As a representative but non-limiting example, useful expression vectors for bacterial use can comprise a selectable marker and bacterial origin of replication derived from commercially available plasmids comprising genetic elements of the well known cloning vector pBR322 (ATCC 37017). Such commercial vectors include, for example, pKK223-3 (Pharmacia Fine Chemicals, Uppsala, Sweden) and GEM 1 (Promega Biotech, Madison, WI, USA). These pBR322 "backbone" sections are combined with an appropriate promoter and the structural sequence to be expressed. Following transformation of a suitable host strain and growth of the host strain to an appropriate cell density, the selected promoter is induced or derepressed by appropriate means (e.g., temperature shift or chemical induction) and cells are cultured for an additional period. Cells are typically harvested by centrifugation, disrupted by physical or chemical means, and the resulting crude extract retained for further purification.

Polynucleotides of the invention can also be used to induce immune responses. For example, as described in Fan et al., *Nat. Biotech.* 17:870-872 (1999), incorporated herein by reference, nucleic acid sequences encoding a polypeptide may be used to generate antibodies against the encoded polypeptide following topical administration of naked plasmid DNA or following injection, and preferably intramuscular injection of the DNA. The nucleic acid sequences are preferably inserted in a recombinant expression vector and may be in the form of naked DNA.

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4.3 ANTISENSE

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Another aspect of the invention pertains to isolated antisense nucleic acid molecules that are hybridizable to or complementary to the nucleic acid molecule comprising the nucleotide sequence of SEQ ID NO: 1-984, 1969-2952, 3937-3942 or 3949-3954, or fragments, analogs or derivatives thereof. An "antisense" nucleic acid comprises a nucleotide sequence that is complementary to a "sense" nucleic acid encoding a protein, *e.g.*, complementary to the coding strand of a double-stranded cDNA molecule or complementary to an mRNA sequence. In specific aspects, antisense nucleic acid molecules are provided that comprise a sequence complementary to at least about 10, 25, 50, 100, 250 or 500 nucleotides or an entire coding strand, or to only a portion thereof. Nucleic acid molecules encoding fragments, homologs, derivatives and analogs of a protein of any of SEQ ID NO: 985-1968, 2953-3936, 3943-3948 or 3955-3960 or antisense nucleic acids complementary to a nucleic acid sequence of SEQ ID NO: 1-984, 1969-2952, 3937-3942 or 3949-3954 are additionally provided.

In one embodiment, an antisense nucleic acid molecule is antisense to a "coding region" of the coding strand of a nucleotide sequence of the invention. The term "coding region" refers to the region of the nucleotide sequence comprising codons which are translated into amino acid residues. In another embodiment, the antisense nucleic acid molecule is antisense to a "noncoding region" of the coding strand of a nucleotide sequence of the invention. The term "noncoding region" refers to 5' and 3' sequences which flank the coding region that are not translated into amino acids (*i.e.*, also referred to as 5' and 3' untranslated regions).

Given the coding strand sequences encoding a nucleic acid disclosed herein (e.g., SEQ ID NO: 1-984, 1969-2952, 3937-3942 or 3949-3954), antisense nucleic acids of the invention can be designed according to the rules of Watson and Crick or Hoogsteen base pairing. The antisense nucleic acid molecule can be complementary to the entire coding region of a mRNA, but more preferably is an oligonucleotide that is antisense to only a portion of the coding or noncoding region of a mRNA. For example, the antisense oligonucleotide can be complementary to the region surrounding the translation start site of a mRNA. An antisense oligonucleotide can be, for example, about 5, 10, 15, 20, 25, 30, 35, 40, 45 or 50 nucleotides in length. An antisense nucleic acid of the invention can be constructed using chemical synthesis or enzymatic ligation reactions using procedures known in the art. For example, an antisense nucleic acid (e.g., an antisense oligonucleotide) can be chemically synthesized using naturally occurring nucleotides or variously modified nucleotides designed to increase the biological stability of the molecules or to increase the physical stability of the duplex formed between the antisense and sense nucleic acids, e.g., phosphorothioate derivatives and acridine substituted nucleotides can be used.

Examples of modified nucleotides that can be used to generate the antisense nucleic acid include: 5-fluorouracil, 5-bromouracil, 5-chlorouracil, 5-iodouracil, hypoxanthine, xanthine, 4-acetylcytosine, 5-(carboxyhydroxylmethyl) uracil, 5-carboxymethylaminomethyl-2-thiouridine, 5-carboxymethylaminomethyluracil, dihydrouracil, beta-D-galactosylqueosine, inosine, N6-isopentenyladenine, 1-methylguanine, 1-methylguanine, 2,2-dimethylguanine, 2-methyladenine, 2-methylguanine, 3-methylcytosine, 5-methylcytosine, N6-adenine, 7-methylguanine, 5-methylaminomethyluracil, 5-methoxyaminomethyl-2-thiouracil, beta-D-mannosylqueosine, 5'-methoxycarboxymethyluracil, 5-methoxyuracil, 2-methylthio-N6-isopentenyladenine, uracil-5-oxyacetic acid (v), wybutoxosine, pseudouracil, queosine, 2-thiocytosine, 5-methyl-2-thiouracil, 2-thiouracil, 4-thiouracil, 5-methyluracil, uracil-5-oxyacetic acid methylester, uracil-5-oxyacetic acid (v), 5-methyl-2-thiouracil, 3-(3-amino-3-N-2-carboxypropyl) uracil, (acp3)w, and 2,6-diaminopurine. Alternatively, the antisense nucleic acid can be produced biologically using an expression vector into which a nucleic acid has been subcloned in an antisense orientation (i.e., RNA transcribed from the inserted nucleic acid will be of an antisense orientation to a target nucleic acid of interest, described further in the following subsection).

The antisense nucleic acid molecules of the invention are typically administered to a subject or generated in situ such that they hybridize with or bind to cellular mRNA and/or genomic DNA encoding a protein according to the invention to thereby inhibit expression of the protein, e.g., by inhibiting transcription and/or translation. The hybridization can be by conventional nucleotide complementarity to form a stable duplex, or, for example, in the case of an antisense nucleic acid molecule that binds to DNA duplexes, through specific interactions in the major groove of the double helix. An example of a route of administration of antisense nucleic acid molecules of the invention includes direct injection at a tissue site. Alternatively, antisense nucleic acid molecules can be modified to target selected cells and then administered systemically. For example, for systemic administration, antisense molecules can be modified such that they specifically bind to receptors or antigens expressed on a selected cell surface, e.g., by linking the antisense nucleic acid molecules to peptides or antibodies that bind to cell surface receptors or antigens. The antisense nucleic acid molecules can also be delivered to cells using the vectors described herein. To achieve sufficient intracellular concentrations of antisense molecules, vector constructs in which the antisense nucleic acid molecule is placed under the control of a strong pol II or pol III promoter are preferred.

In yet another embodiment, the antisense nucleic acid molecule of the invention is an α -anomeric nucleic acid molecule. An α -anomeric nucleic acid molecule forms specific double-stranded hybrids with complementary RNA in which, contrary to the usual β -units, the

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strands run parallel to each other (Gaultier et al. (1987) Nucleic Acids Res 15: 6625-6641). The antisense nucleic acid molecule can also comprise a 2'-o-methylribonucleotide (Inoue et al. (1987) Nucleic Acids Res 15: 6131-6148) or a chimeric RNA -DNA analogue (Inoue et al. (1987) FEBS Lett 215: 327-330).

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4.4 RIBOZYMES AND PNA MOIETIES

In still another embodiment, an antisense nucleic acid of the invention is a ribozyme. Ribozymes are catalytic RNA molecules with ribonuclease activity that are capable of cleaving a single-stranded nucleic acid, such as a mRNA, to which they have a complementary region. Thus, ribozymes (e.g., hammerhead ribozymes (described in Haselhoff and Gerlach (1988) Nature 334:585-591)) can be used to catalytically cleave a mRNA transcripts to thereby inhibit translation of a mRNA. A ribozyme having specificity for a nucleic acid of the invention can be designed based upon the nucleotide sequence of a DNA disclosed herein (i.e., SEQ ID NO: 1-984, 1969-2952, 3937-3942 or 3949-3954). For example, a derivative of a Tetrahymena L-19 IVS RNA can be constructed in which the nucleotide sequence of the active site is complementary to the nucleotide sequence to be cleaved in a SECX-encoding mRNA. See, e.g., Cech et al. U.S. Pat. No. 4,987,071; and Cech et al. U.S. Pat. No. 5,116,742. Alternatively, SECX mRNA can be used to select a catalytic RNA having a specific ribonuclease activity from a pool of RNA molecules. See, e.g., Bartel et al., (1993) Science 261:1411-1418.

Alternatively, gene expression can be inhibited by targeting nucleotide sequences complementary to the regulatory region (e.g., promoter and/or enhancers) to form triple helical structures that prevent transcription of the gene in target cells. See generally, Helene. (1991)

Anticancer Drug Des. 6: 569-84; Helene. et al. (1992) Ann. N.Y. Acad. Sci. 660:27-36; and Maher (1992) Bioassays 14: 807-15.

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In various embodiments, the nucleic acids of the invention can be modified at the base moiety, sugar moiety or phosphate backbone to improve, e.g., the stability, hybridization, or solubility of the molecule. For example, the deoxyribose phosphate backbone of the nucleic acids can be modified to generate peptide nucleic acids (see Hyrup et al. (1996) Bioorg Med Chem 4: 5-23). As used herein, the terms "peptide nucleic acids" or "PNAs" refer to nucleic acid mimics, e.g., DNA mimics, in which the deoxyribose phosphate backbone is replaced by a pseudopeptide backbone and only the four natural nucleobases are retained. The neutral backbone of PNAs has been shown to allow for specific hybridization to DNA and RNA under conditions of low ionic strength. The synthesis of PNA oligomers can be performed using standard solid phase peptide synthesis protocols as described in Hyrup et al. (1996) above; Perry-O'Keefe et al. (1996) PNAS 93: 14670-675.

PNAs of the invention can be used in therapeutic and diagnostic applications. For example, PNAs can be used as antisense or antigene agents for sequence-specific modulation of gene expression by, e.g., inducing transcription or translation arrest or inhibiting replication. PNAs of the invention can also be used, e.g., in the analysis of single base pair mutations in a gene by, e.g., PNA directed PCR clamping; as artificial restriction enzymes when used in combination with other enzymes, e.g., S1 nucleases (Hyrup B. (1996) above); or as probes or primers for DNA sequence and hybridization (Hyrup et al. (1996), above; Perry-O'Keefe (1996), above).

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In another embodiment, PNAs of the invention can be modified, e.g., to enhance their stability or cellular uptake, by attaching lipophilic or other helper groups to PNA, by the formation of PNA-DNA chimeras, or by the use of liposomes or other techniques of drug delivery known in the art. For example, PNA-DNA chimeras can be generated that may combine the advantageous properties of PNA and DNA. Such chimeras allow DNA recognition enzymes, e.g., RNase H and DNA polymerases, to interact with the DNA portion while the PNA portion would provide high binding affinity and specificity. PNA-DNA chimeras can be linked using linkers of appropriate lengths selected in terms of base stacking, number of bonds between the nucleobases, and orientation (Hyrup (1996) above). The synthesis of PNA-DNA chimeras can be performed as described in Hyrup (1996) above and Finn et al. (1996) Nucl Acids Res 24: 3357-63. For example, a DNA chain can be synthesized on a solid support using standard phosphoramidite coupling chemistry, and modified nucleoside analogs, e.g., 5'-(4-methoxytrityl)amino-5'-deoxy-thymidine phosphoramidite, can be used between the PNA and the 5' end of DNA (Mag et al. (1989) Nucl Acid Res 17: 5973-88). PNA monomers are then coupled in a stepwise manner to produce a chimeric molecule with a 5' PNA segment and a 3' DNA segment (Finn et al. (1996) above). Alternatively, chimeric molecules can be synthesized with a 5' DNA segment and a 3' PNA segment. See, Petersen et al. (1975) Bioorg Med Chem Lett 5: 1119-11124.

In other embodiments, the oligonucleotide may include other appended groups such as peptides (e.g., for targeting host cell receptors in vivo), or agents facilitating transport across the cell membrane (see, e.g., Letsinger et al., 1989, Proc. Natl. Acad. Sci. U.S.A. 86:6553-6556; Lemaitre et al., 1987, Proc. Natl. Acad. Sci. 84:648-652; PCT Publication No. W088/09810) or the blood-brain barrier (see, e.g., PCT Publication No. W089/10134). In addition, oligonucleotides can be modified with hybridization triggered cleavage agents (See, e.g., Krol et al., 1988, BioTechniques 6:958-976) or intercalating agents. (See, e.g., Zon, 1988, Pharm. Res. 5: 539-549). To this end, the oligonucleotide may be conjugated to another molecule, e.g., a

peptide, a hybridization triggered cross-linking agent, a transport agent, a hybridization-triggered cleavage agent, etc.

4.5 HOSTS

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The present invention further provides host cells genetically engineered to contain the polynucleotides of the invention. For example, such host cells may contain nucleic acids of the invention introduced into the host cell using known transformation, transfection or infection methods. The present invention still further provides host cells genetically engineered to express the polynucleotides of the invention, wherein such polynucleotides are in operative association with a regulatory sequence heterologous to the host cell which drives expression of the polynucleotides in the cell.

Knowledge of nucleic acid sequences allows for modification of cells to permit, or increase, expression of endogenous polypeptide. Cells can be modified (e.g., by homologous recombination) to provide increased polypeptide expression by replacing, in whole or in part, the naturally occurring promoter with all or part of a heterologous promoter so that the cells express the polypeptide at higher levels. The heterologous promoter is inserted in such a manner that it is operatively linked to the encoding sequences. See, for example, PCT International Publication No. WO94/12650, PCT International Publication No. WO92/20808, and PCT International Publication No. WO91/09955. It is also contemplated that, in addition to heterologous promoter DNA, amplifiable marker DNA (e.g., ada, dhfr, and the multifunctional CAD gene which encodes carbamyl phosphate synthase, aspartate transcarbamylase, and dihydroorotase) and/or intron DNA may be inserted along with the heterologous promoter DNA. If linked to the coding sequence, amplification of the marker DNA by standard selection methods results in coamplification of the desired protein coding sequences in the cells.

The host cell can be a higher eukaryotic host cell, such as a mammalian cell, a lower eukaryotic host cell, such as a yeast cell, or the host cell can be a prokaryotic cell, such as a bacterial cell. Introduction of the recombinant construct into the host cell can be effected by calcium phosphate transfection, DEAE, dextran mediated transfection, or electroporation (Davis, L. et al., *Basic Methods in Molecular Biology* (1986)). The host cells containing one of the polynucleotides of the invention, can be used in conventional manners to produce the gene product encoded by the isolated fragment (in the case of an ORF) or can be used to produce a heterologous protein under the control of the EMF.

Any host/vector system can be used to express one or more of the ORFs of the present invention. These include, but are not limited to, eukaryotic hosts such as HeLa cells, Cv-1 cell, COS cells, 293 cells, and Sf9 cells, as well as prokaryotic host such as *E. coli* and *B. subtilis*.

The most preferred cells are those which do not normally express the particular polypeptide or protein or which expresses the polypeptide or protein at low natural level. Mature proteins can be expressed in mammalian cells, yeast, bacteria, or other cells under the control of appropriate promoters. Cell-free translation systems can also be employed to produce such proteins using RNAs derived from the DNA constructs of the present invention. Appropriate cloning and expression vectors for use with prokaryotic and eukaryotic hosts are described by Sambrook, et al., in Molecular Cloning: A Laboratory Manual, Second Edition, Cold Spring Harbor, New York (1989), the disclosure of which is hereby incorporated by reference.

Various mammalian cell culture systems can also be employed to express recombinant protein. Examples of mammalian expression systems include the COS-7 lines of monkey kidney fibroblasts, described by Gluzman, Cell 23:175 (1981). Other cell lines capable of expressing a compatible vector are, for example, the C127, monkey COS cells, Chinese Hamster Ovary (CHO) cells, human kidney 293 cells, human epidermal A431 cells, human Colo205 cells, 3T3 cells, CV-1 cells, other transformed primate cell lines, normal diploid cells, cell strains derived from in vitro culture of primary tissue, primary explants, HeLa cells, mouse L cells, BHK, HL-60, U937, HaK or Jurkat cells. Mammalian expression vectors will comprise an origin of replication, a suitable promoter and also any necessary ribosome binding sites, polyadenylation site, splice donor and acceptor sites, transcriptional termination sequences, and 5' flanking nontranscribed sequences. DNA sequences derived from the SV40 viral genome, for example, SV40 origin, early promoter, enhancer, splice, and polyadenylation sites may be used to provide the required nontranscribed genetic elements. Recombinant polypeptides and proteins produced in bacterial culture are usually isolated by initial extraction from cell pellets, followed by one or more salting-out, aqueous ion exchange or size exclusion chromatography steps. Protein refolding steps can be used, as necessary, in completing configuration of the mature protein. Finally, high performance liquid chromatography (HPLC) can be employed for final purification steps. Microbial cells employed in expression of proteins can be disrupted by any convenient method, including freeze-thaw cycling, sonication, mechanical disruption, or use of cell lysing agents.

Alternatively, it may be possible to produce the protein in lower eukaryotes such as yeast or insects or in prokaryotes such as bacteria. Potentially suitable yeast strains include Saccharomyces cerevisiae, Schizosaccharomyces pombe, Kluyveromyces strains, Candida, or any yeast strain capable of expressing heterologous proteins. Potentially suitable bacterial strains include Escherichia coli, Bacillus subtilis, Salmonella typhimurium, or any bacterial strain capable of expressing heterologous proteins. If the protein is made in yeast or bacteria, it may be necessary to modify the protein produced therein, for example by phosphorylation or

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glycosylation of the appropriate sites, in order to obtain the functional protein. Such covalent attachments may be accomplished using known chemical or enzymatic methods.

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In another embodiment of the present invention, cells and tissues may be engineered to express an endogenous gene comprising the polynucleotides of the invention under the control of inducible regulatory elements, in which case the regulatory sequences of the endogenous gene may be replaced by homologous recombination. As described herein, gene targeting can be used to replace a gene's existing regulatory region with a regulatory sequence isolated from a different gene or a novel regulatory sequence synthesized by genetic engineering methods. Such regulatory sequences may be comprised of promoters, enhancers, scaffold-attachment regions, negative regulatory elements, transcriptional initiation sites, regulatory protein binding sites or combinations of said sequences. Alternatively, sequences which affect the structure or stability of the RNA or protein produced may be replaced, removed, added, or otherwise modified by targeting. These sequence include polyadenylation signals, mRNA stability elements, splice sites, leader sequences for enhancing or modifying transport or secretion properties of the protein, or other sequences which alter or improve the function or stability of protein or RNA molecules.

The targeting event may be a simple insertion of the regulatory sequence, placing the gene under the control of the new regulatory sequence, e.g., inserting a new promoter or enhancer or both upstream of a gene. Alternatively, the targeting event may be a simple deletion of a regulatory element, such as the deletion of a tissue-specific negative regulatory element. Alternatively, the targeting event may replace an existing element; for example, a tissue-specific enhancer can be replaced by an enhancer that has broader or different cell-type specificity than the naturally occurring elements. Here, the naturally occurring sequences are deleted and new sequences are added. In all cases, the identification of the targeting event may be facilitated by the use of one or more selectable marker genes that are contiguous with the targeting DNA, allowing for the selection of cells in which the exogenous DNA has integrated into the host cell genome. The identification of the targeting event may also be facilitated by the use of one or more marker genes exhibiting the property of negative selection, such that the negatively selectable marker is linked to the exogenous DNA, but configured such that the negatively selectable marker flanks the targeting sequence, and such that a correct homologous recombination event with sequences in the host cell genome does not result in the stable integration of the negatively selectable marker. Markers useful for this purpose include the Herpes Simplex Virus thymidine kinase (TK) gene or the bacterial xanthine-guanine phosphoribosyl-transferase (gpt) gene.

The gene targeting or gene activation techniques which can be used in accordance with this aspect of the invention are more particularly described in U.S. Patent No. 5,272,071 to Chappel; U.S. Patent No. 5,578,461 to Sherwin et al.; International Application No. PCT/US92/09627 (WO93/09222) by Selden et al.; and International Application No. PCT/US90/06436 (WO91/06667) by Skoultchi et al., each of which is incorporated by reference herein in its entirety.

4.6 POLYPEPTIDES OF THE INVENTION

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The isolated polypeptides of the invention include, but are not limited to, a polypeptide comprising: the amino acid sequences set forth as any one of SEQ ID NO: 985-1968, 2953-3936, 3943-3948 or 3955-3960 or an amino acid sequence encoded by any one of the nucleotide sequences SEQ ID NO: 1-984, 1969-2952, 3937-3942 or 3949-3954 or the corresponding full length or mature protein. Polypeptides of the invention also include polypeptides preferably with biological or immunological activity that are encoded by: (a) a polynucleotide having any one of the nucleotide sequences set forth in SEQ ID NO: 1-984, 1969-2952, 3937-3942 or 3949-3954 or (b) polynucleotides encoding any one of the amino acid sequences set forth as SEO ID NO: 985-1968, 2953-3936, 3943-3948 or 3955-3960 or (c) polynucleotides that hybridize to the complement of the polynucleotides of either (a) or (b) under stringent hybridization conditions. The invention also provides biologically active or immunologically active variants of any of the amino acid sequences set forth as SEQ ID NO: 985-1968, 2953-3936, 3943-3948 or 3955-3960 or the corresponding full length or mature protein; and "substantial equivalents" thereof (e.g., at least about 65%, at least about 70%, at least about 75%, at least about 80%, 81%, 82%, 83%, 84%, more typically at least about 85%, 86%, 87%, 88%, 89%, and more typically at least about 90%, 91%, 92%, 93%, 94%, and even more typically at least about 95%, 96%, 97%, 98%, 99%. sequence identity that retain biological activity. Polypeptides encoded by allelic variants may have a similar, increased, or decreased activity compared to polypeptides comprising SEO ID NO: 985-1968, 2953-3936, 3943-3948 or 3955-3960.

Fragments of the proteins of the present invention which are capable of exhibiting biological activity are also encompassed by the present invention. Fragments of the protein may be in linear form or they may be cyclized using known methods, for example, as described in H. U. Saragovi, et al., Bio/Technology 10, 773-778 (1992) and in R. S. McDowell, et al., J. Amer. Chem. Soc. 114, 9245-9253 (1992), both of which are incorporated herein by reference. Such fragments may be fused to carrier molecules such as immunoglobulins for many purposes, including increasing the valency of protein binding sites.

The present invention also provides both full-length and mature forms (for example, without a signal sequence or precursor sequence) of the disclosed proteins. The protein coding sequence is identified in the sequence listing by translation of the disclosed nucleotide sequences. The mature form of such protein may be obtained by expression of a full-length polynucleotide in a suitable mammalian cell or other host cell. The sequence of the mature form of the protein is also determinable from the amino acid sequence of the full-length form. Where proteins of the present invention are membrane bound, soluble forms of the proteins are also provided. In such forms, part or all of the regions causing the proteins to be membrane bound are deleted so that the proteins are fully secreted from the cell in which they are expressed.

Protein compositions of the present invention may further comprise an acceptable carrier, such as a hydrophilic, e.g., pharmaceutically acceptable, carrier.

The present invention further provides isolated polypeptides encoded by the nucleic acid fragments of the present invention or by degenerate variants of the nucleic acid fragments of the present invention. By "degenerate variant" is intended nucleotide fragments which differ from a nucleic acid fragment of the present invention (e.g., an ORF) by nucleotide sequence but, due to the degeneracy of the genetic code, encode an identical polypeptide sequence. Preferred nucleic acid fragments of the present invention are the ORFs that encode proteins.

A variety of methodologies known in the art can be utilized to obtain any one of the isolated polypeptides or proteins of the present invention. At the simplest level, the amino acid sequence can be synthesized using commercially available peptide synthesizers. The synthetically-constructed protein sequences, by virtue of sharing primary, secondary or tertiary structural and/or conformational characteristics with proteins may possess biological properties in common therewith, including protein activity. This technique is particularly useful in producing small peptides and fragments of larger polypeptides. Fragments are useful, for example, in generating antibodies against the native polypeptide. Thus, they may be employed as biologically active or immunological substitutes for natural, purified proteins in screening of therapeutic compounds and in immunological processes for the development of antibodies.

The polypeptides and proteins of the present invention can alternatively be purified from cells which have been altered to express the desired polypeptide or protein. As used herein, a cell is said to be altered to express a desired polypeptide or protein when the cell, through genetic manipulation, is made to produce a polypeptide or protein which it normally does not produce or which the cell normally produces at a lower level. One skilled in the art can readily adapt procedures for introducing and expressing either recombinant or synthetic sequences into eukaryotic or prokaryotic cells in order to generate a cell which produces one of the polypeptides or proteins of the present invention.

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The invention also relates to methods for producing a polypeptide comprising growing a culture of host cells of the invention in a suitable culture medium, and purifying the protein from the cells or the culture in which the cells are grown. For example, the methods of the invention include a process for producing a polypeptide in which a host cell containing a suitable expression vector that includes a polynucleotide of the invention is cultured under conditions that allow expression of the encoded polypeptide. The polypeptide can be recovered from the culture, conveniently from the culture medium, or from a lysate prepared from the host cells and further purified. Preferred embodiments include those in which the protein produced by such process is a full length or mature form of the protein.

In an alternative method, the polypeptide or protein is purified from bacterial cells which naturally produce the polypeptide or protein. One skilled in the art can readily follow known methods for isolating polypeptides and proteins in order to obtain one of the isolated polypeptides or proteins of the present invention. These include, but are not limited to, immunochromatography, HPLC, size-exclusion chromatography, ion-exchange chromatography, and immuno-affinity chromatography. See, e.g., Scopes, Protein Purification: Principles and Practice, Springer-Verlag (1994); Sambrook, et al., in Molecular Cloning: A Laboratory Manual; Ausubel et al., Current Protocols in Molecular Biology. Polypeptide fragments that retain biological/immunological activity include fragments comprising greater than about 100 amino acids, or greater than about 200 amino acids, and fragments that encode specific protein domains.

The purified polypeptides can be used in *in vitro* binding assays which are well known in the art to identify molecules which bind to the polypeptides. These molecules include but are not limited to, for e.g., small molecules, molecules from combinatorial libraries, antibodies or other proteins. The molecules identified in the binding assay are then tested for antagonist or agonist activity in *in vivo* tissue culture or animal models that are well known in the art. In brief, the molecules are titrated into a plurality of cell cultures or animals and then tested for either cell/animal death or prolonged survival of the animal/cells.

In addition, the peptides of the invention or molecules capable of binding to the peptides may be complexed with toxins, e.g., ricin or cholera, or with other compounds that are toxic to cells. The toxin-binding molecule complex is then targeted to a tumor or other cell by the specificity of the binding molecule for SEQ ID NO: 985-1968, 2953-3936, 3943-3948 or 3955-3960.

The protein of the invention may also be expressed as a product of transgenic animals, e.g., as a component of the milk of transgenic cows, goats, pigs, or sheep which are characterized by somatic or germ cells containing a nucleotide sequence encoding the protein.

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The proteins provided herein also include proteins characterized by amino acid sequences similar to those of purified proteins but into which modification are naturally provided or deliberately engineered. For example, modifications, in the peptide or DNA sequence, can be made by those skilled in the art using known techniques. Modifications of interest in the protein sequences may include the alteration, substitution, replacement, insertion or deletion of a selected amino acid residue in the coding sequence. For example, one or more of the cysteine residues may be deleted or replaced with another amino acid to alter the conformation of the molecule. Techniques for such alteration, substitution, replacement, insertion or deletion are well known to those skilled in the art (see, e.g., U.S. Pat. No. 4,518,584). Preferably, such alteration, substitution, replacement, insertion or deletion retains the desired activity of the protein. Regions of the protein that are important for the protein function can be determined by various methods known in the art including the alanine-scanning method which involved systematic substitution of single or strings of amino acids with alanine, followed by testing the resulting alanine-containing variant for biological activity. This type of analysis determines the importance of the substituted amino acid(s) in biological activity. Regions of the protein that are important for protein function may be determined by the eMATRIX program.

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Other fragments and derivatives of the sequences of proteins which would be expected to retain protein activity in whole or in part and are useful for screening or other immunological methodologies may also be easily made by those skilled in the art given the disclosures herein. Such modifications are encompassed by the present invention.

The protein may also be produced by operably linking the isolated polynucleotide of the invention to suitable control sequences in one or more insect expression vectors, and employing an insect expression system. Materials and methods for baculovirus/insect cell expression systems are commercially available in kit form from, e.g., Invitrogen, San Diego, Calif., U.S.A. (the MaxBatTM kit), and such methods are well known in the art, as described in Summers and Smith, Texas Agricultural Experiment Station Bulletin No. 1555 (1987), incorporated herein by reference. As used herein, an insect cell capable of expressing a polynucleotide of the present invention is "transformed."

The protein of the invention may be prepared by culturing transformed host cells under culture conditions suitable to express the recombinant protein. The resulting expressed protein may then be purified from such culture (i.e., from culture medium or cell extracts) using known purification processes, such as gel filtration and ion exchange chromatography. The purification of the protein may also include an affinity column containing agents which will bind to the protein; one or more column steps over such affinity resins as concanavalin A-agarose, heparin-toyopearlTM or Cibacrom blue 3GA SepharoseTM; one or more steps involving

hydrophobic interaction chromatography using such resins as phenyl ether, butyl ether, or propyl ether; or immunoaffinity chromatography.

Alternatively, the protein of the invention may also be expressed in a form which will facilitate purification. For example, it may be expressed as a fusion protein, such as those of maltose binding protein (MBP), glutathione-S-transferase (GST) or thioredoxin (TRX), or as a His tag. Kits for expression and purification of such fusion proteins are commercially available from New England BioLab (Beverly, Mass.), Pharmacia (Piscataway, N.J.) and Invitrogen, respectively. The protein can also be tagged with an epitope and subsequently purified by using a specific antibody directed to such epitope. One such epitope ("FLAG®") is commercially available from Kodak (New Haven, Conn.).

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Finally, one or more reverse-phase high performance liquid chromatography (RP-HPLC) steps employing hydrophobic RP-HPLC media, e.g., silica gel having pendant methyl or other aliphatic groups, can be employed to further purify the protein. Some or all of the foregoing purification steps, in various combinations, can also be employed to provide a substantially homogeneous isolated recombinant protein. The protein thus purified is substantially free of other mammalian proteins and is defined in accordance with the present invention as an "isolated protein."

The polypeptides of the invention include analogs (variants). This embraces fragments, as well as peptides in which one or more amino acids has been deleted, inserted, or substituted. Also, analogs of the polypeptides of the invention embrace fusions of the polypeptides or modifications of the polypeptides of the invention, wherein the polypeptide or analog is fused to another moiety or moieties, e.g., targeting moiety or another therapeutic agent. Such analogs may exhibit improved properties such as activity and/or stability. Examples of moieties which may be fused to the polypeptide or an analog include, for example, targeting moieties which provide for the delivery of polypeptide to pancreatic cells, e.g., antibodies to pancreatic cells, antibodies to immune cells such as T-cells, monocytes, dendritic cells, granulocytes, etc., as well as receptor and ligands expressed on pancreatic or immune cells. Other moieties which may be fused to the polypeptide include therapeutic agents which are used for treatment, for example, immunosuppressive drugs such as cyclosporin, SK506, azathioprine, CD3 antibodies and steroids. Also, polypeptides may be fused to immune modulators, and other cytokines such as alpha or beta interferon.

4.6.1 DETERMINING POLYPEPTIDE AND POLYNUCLEOTIDE IDENTITY AND SIMILARITY

Preferred identity and/or similarity are designed to give the largest match between the sequences tested. Methods to determine identity and similarity are codified in computer programs including, but are not limited to, the GCG program package, including GAP (Devereux, J., et al., Nucleic Acids Research 12(1):387 (1984); Genetics Computer Group, University of Wisconsin, Madison, WI), BLASTP, BLASTN, BLASTX, FASTA (Altschul, S.F. et al., J. Molec. Biol. 215:403-410 (1990), PSI-BLAST (Altschul S.F. et al., Nucleic Acids Res. vol. 25, pp. 3389-3402, herein incorporated by reference), eMatrix software (Wu et al., J. Comp. Biol., Vol. 6, pp. 219-235 (1999), herein incorporated by reference), eMotif software (Nevill-Manning et al, ISMB-97, Vol. 4, pp. 202-209, herein incorporated by reference), pFam software (Sonnhammer et al., Nucleic Acids Res., Vol. 26(1), pp. 320-322 (1998), herein incorporated by reference) and the Kyte-Doolittle hydrophobocity prediction algorithm (J. Mol Biol, 157, pp. 105-31 (1982), incorporated herein by reference). The BLAST programs are publicly available from the National Center for Biotechnology Information (NCBI) and other sources (BLAST Manual, Altschul, S., et al. NCB NLM NIH Bethesda, MD 20894; Altschul, S., et al., J. Mol. Biol. 215:403-410 (1990).

4.7 CHIMERIC AND FUSION PROTEINS

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The invention also provides chimeric or fusion proteins. As used herein, a "chimeric protein" or "fusion protein" comprises a polypeptide of the invention operatively linked to another polypeptide. Within a fusion protein the polypeptide according to the invention can correspond to all or a portion of a protein according to the invention. In one embodiment, a fusion protein comprises at least one biologically active portion of a protein according to the invention. In another embodiment, a fusion protein comprises at least two biologically active portions of a protein according to the invention. Within the fusion protein, the term "operatively linked" is intended to indicate that the polypeptide according to the invention and the other polypeptide are fused in-frame to each other. The polypeptide can be fused to the N-terminus or C-terminus.

For example, in one embodiment a fusion protein comprises a polypeptide according to the invention operably linked to the extracellular domain of a second protein.

In another embodiment, the fusion protein is a GST-fusion protein in which the polypeptide sequences of the invention are fused to the C-terminus of the GST (i.e., glutathione S-transferase) sequences.

In another embodiment, the fusion protein is an immunoglobulin fusion protein in which the polypeptide sequences according to the invention comprise one or more domains fused to sequences derived from a member of the immunoglobulin protein family. The immunoglobulin fusion proteins of the invention can be incorporated into pharmaceutical compositions and ., 0 0210.20

administered to a subject to inhibit an interaction between a ligand and a protein of the invention on the surface of a cell, to thereby suppress signal transduction *in vivo*. The immunoglobulin fusion proteins can be used to affect the bioavailability of a cognate ligand. Inhibition of the ligand/protein interaction may be useful therapeutically for both the treatment of proliferative and differentiative disorders, *e,g.*, cancer as well as modulating (*e.g.*, promoting or inhibiting) cell survival. Moreover, the immunoglobulin fusion proteins of the invention can be used as immunogens to produce antibodies in a subject, to purify ligands, and in screening assays to identify molecules that inhibit the interaction of a polypeptide of the invention with a ligand.

A chimeric or fusion protein of the invention can be produced by standard recombinant DNA techniques. For example, DNA fragments coding for the different polypeptide sequences are ligated together in-frame in accordance with conventional techniques, e.g., by employing blunt-ended or stagger-ended termini for ligation, restriction enzyme digestion to provide for appropriate termini, filling-in of cohesive ends as appropriate, alkaline phosphatase treatment to avoid undesirable joining, and enzymatic ligation. In another embodiment, the fusion gene can be synthesized by conventional techniques including automated DNA synthesizers.

Alternatively, PCR amplification of gene fragments can be carried out using anchor primers that give rise to complementary overhangs between two consecutive gene fragments that can subsequently be annealed and reamplified to generate a chimeric gene sequence (see, for example, Ausubel et al. (eds.) Current Protocols in Molecular Biology, John Wiley & Sons, 1992). Moreover, many expression vectors are commercially available that already encode a fusion moiety (e.g., a GST polypeptide). A nucleic acid encoding a polypeptide of the invention can be cloned into such an expression vector such that the fusion moiety is linked in-frame to the protein of the invention.

25 4.8 GENE THERAPY

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Mutations in the polynucleotides of the invention gene may result in loss of normal function of the encoded protein. The invention thus provides gene therapy to restore normal activity of the polypeptides of the invention; or to treat disease states involving polypeptides of the invention. Delivery of a functional gene encoding polypeptides of the invention to appropriate cells is effected *ex vivo*, *in situ*, or *in vivo* by use of vectors, and more particularly viral vectors (e.g., adenovirus, adeno-associated virus, or a retrovirus), or *ex vivo* by use of physical DNA transfer methods (e.g., liposomes or chemical treatments). See, for example, Anderson, Nature, supplement to vol. 392, no. 6679, pp.25-20 (1998). For additional reviews of gene therapy technology see Friedmann, Science, 244: 1275-1281 (1989); Verma, Scientific American: 68-84 (1990); and Miller, Nature, 357: 455-460 (1992). Introduction of any one of

the nucleotides of the present invention or a gene encoding the polypeptides of the present invention can also be accomplished with extrachromosomal substrates (transient expression) or artificial chromosomes (stable expression). Cells may also be cultured *ex vivo* in the presence of proteins of the present invention in order to proliferate or to produce a desired effect on or activity in such cells. Treated cells can then be introduced *in vivo* for therapeutic purposes. Alternatively, it is contemplated that in other human disease states, preventing the expression of or inhibiting the activity of polypeptides of the invention will be useful in treating the disease states. It is contemplated that antisense therapy or gene therapy could be applied to negatively regulate the expression of polypeptides of the invention.

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Other methods inhibiting expression of a protein include the introduction of antisense molecules to the nucleic acids of the present invention, their complements, or their translated RNA sequences, by methods known in the art. Further, the polypeptides of the present invention can be inhibited by using targeted deletion methods, or the insertion of a negative regulatory element such as a silencer, which is tissue specific.

The present invention still further provides cells genetically engineered *in vivo* to express the polynucleotides of the invention, wherein such polynucleotides are in operative association with a regulatory sequence heterologous to the host cell which drives expression of the polynucleotides in the cell. These methods can be used to increase or decrease the expression of the polynucleotides of the present invention.

Knowledge of DNA sequences provided by the invention allows for modification of cells to permit, increase, or decrease, expression of endogenous polypeptide. Cells can be modified (e.g., by homologous recombination) to provide increased polypeptide expression by replacing, in whole or in part, the naturally occurring promoter with all or part of a heterologous promoter so that the cells express the protein at higher levels. The heterologous promoter is inserted in such a manner that it is operatively linked to the desired protein encoding sequences. See, for example, PCT International Publication No. WO 94/12650, PCT International Publication No. WO 92/20808, and PCT International Publication No. WO 91/09955. It is also contemplated that, in addition to heterologous promoter DNA, amplifiable marker DNA (e.g., ada, dhfr, and the multifunctional CAD gene which encodes carbamyl phosphate synthase, aspartate transcarbamylase, and dihydroorotase) and/or intron DNA may be inserted along with the heterologous promoter DNA. If linked to the desired protein coding sequence, amplification of the marker DNA by standard selection methods results in co-amplification of the desired protein coding sequences in the cells.

In another embodiment of the present invention, cells and tissues may be engineered to express an endogenous gene comprising the polynucleotides of the invention under the control of inducible regulatory elements, in which case the regulatory sequences of the endogenous gene may

be replaced by homologous recombination. As described herein, gene targeting can be used to replace a gene's existing regulatory region with a regulatory sequence isolated from a different gene or a novel regulatory sequence synthesized by genetic engineering methods. Such regulatory sequences may be comprised of promoters, enhancers, scaffold-attachment regions, negative regulatory elements, transcriptional initiation sites, regulatory protein binding sites or combinations of said sequences. Alternatively, sequences which affect the structure or stability of the RNA or protein produced may be replaced, removed, added, or otherwise modified by targeting. These sequences include polyadenylation signals, mRNA stability elements, splice sites, leader sequences for enhancing or modifying transport or secretion properties of the protein, or other sequences which alter or improve the function or stability of protein or RNA molecules.

The targeting event may be a simple insertion of the regulatory sequence, placing the gene under the control of the new regulatory sequence, e.g., inserting a new promoter or enhancer or both upstream of a gene. Alternatively, the targeting event may be a simple deletion of a regulatory element, such as the deletion of a tissue-specific negative regulatory element. Alternatively, the targeting event may replace an existing element; for example, a tissue-specific enhancer can be replaced by an enhancer that has broader or different cell-type specificity than the naturally occurring elements. Here, the naturally occurring sequences are deleted and new sequences are added. In all cases, the identification of the targeting event may be facilitated by the use of one or more selectable marker genes that are contiguous with the targeting DNA, allowing for the selection of cells in which the exogenous DNA has integrated into the cell genome. The identification of the targeting event may also be facilitated by the use of one or more marker genes exhibiting the property of negative selection, such that the negatively selectable marker is linked to the exogenous DNA, but configured such that the negatively selectable marker flanks the targeting sequence, and such that a correct homologous recombination event with sequences in the host cell genome does not result in the stable integration of the negatively selectable marker. Markers useful for this purpose include the Herpes Simplex Virus thymidine kinase (TK) gene or the bacterial xanthine-guanine phosphoribosyl-transferase (gpt) gene.

The gene targeting or gene activation techniques which can be used in accordance with this aspect of the invention are more particularly described in U.S. Patent No. 5,272,071 to Chappel; U.S. Patent No. 5,578,461 to Sherwin et al.; International Application No. PCT/US92/09627 (WO93/09222) by Selden et al.; and International Application No. PCT/US90/06436 (WO91/06667) by Skoultchi et al., each of which is incorporated by reference herein in its entirety.

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In preferred methods to determine biological functions of the polypeptides of the invention in vivo, one or more genes provided by the invention are either over expressed or inactivated in the germ line of animals using homologous recombination [Capecchi, Science 244:1288-1292 (1989)]. Animals in which the gene is over expressed, under the regulatory control of exogenous or endogenous promoter elements, are known as transgenic animals. Animals in which an endogenous gene has been inactivated by homologous recombination are referred to as "knockout" animals. Knockout animals, preferably non-human mammals, can be prepared as described in U.S. Patent No. 5,557,032, incorporated herein by reference. Transgenic animals are useful to determine the roles polypeptides of the invention play in biological processes, and preferably in disease states. Transgenic animals are useful as model systems to identify compounds that modulate lipid metabolism. Transgenic animals, preferably non-human mammals, are produced using methods as described in U.S. Patent No 5,489,743 and PCT Publication No. WO94/28122, incorporated herein by reference.

Transgenic animals can be prepared wherein all or part of a promoter of the polynucleotides of the invention is either activated or inactivated to alter the level of expression of the polypeptides of the invention. Inactivation can be carried out using homologous recombination methods described above. Activation can be achieved by supplementing or even replacing the homologous promoter to provide for increased protein expression. The homologous promoter can be supplemented by insertion of one or more heterologous enhancer elements known to confer promoter activation in a particular tissue.

The polynucleotides of the present invention also make possible the development, through, e.g., homologous recombination or knock out strategies, of animals that fail to express polypeptides of the invention or that express a variant polypeptide. Such animals are useful as models for studying the *in vivo* activities of polypeptide as well as for studying modulators of the polypeptides of the invention.

In preferred methods to determine biological functions of the polypeptides of the invention *in vivo*, one or more genes provided by the invention are either over expressed or inactivated in the germ line of animals using homologous recombination [Capecchi, Science 244:1288-1292 (1989)]. Animals in which the gene is over expressed, under the regulatory control of exogenous or endogenous promoter elements, are known as transgenic animals. Animals in which an endogenous gene has been inactivated by homologous recombination are referred to as "knockout" animals. Knockout animals, preferably non-human mammals, can be prepared as described in U.S. Patent No. 5,557,032, incorporated herein by reference. Transgenic animals are useful to determine the roles polypeptides of the invention play in biological processes, and preferably in disease states. Transgenic animals are useful as model systems to

identify compounds that modulate lipid metabolism. Transgenic animals, preferably non-human mammals, are produced using methods as described in U.S. Patent No 5,489,743 and PCT Publication No. WO94/28122, incorporated herein by reference.

Transgenic animals can be prepared wherein all or part of the polynucleotides of the invention promoter is either activated or inactivated to alter the level of expression of the polypeptides of the invention. Inactivation can be carried out using homologous recombination methods described above. Activation can be achieved by supplementing or even replacing the homologous promoter to provide for increased protein expression. The homologous promoter can be supplemented by insertion of one or more heterologous enhancer elements known to confer promoter activation in a particular tissue.

4.10 USES AND BIOLOGICAL ACTIVITY

The polynucleotides and proteins of the present invention are expected to exhibit one or more of the uses or biological activities (including those associated with assays cited herein) identified herein. Uses or activities described for proteins of the present invention may be provided by administration or use of such proteins or of polynucleotides encoding such proteins (such as, for example, in gene therapies or vectors suitable for introduction of DNA). The mechanism underlying the particular condition or pathology will dictate whether the polypeptides of the invention, the polynucleotides of the invention or modulators (activators or inhibitors) thereof would be beneficial to the subject in need of treatment. Thus, "therapeutic compositions of the invention" include compositions comprising isolated polynucleotides (including recombinant DNA molecules, cloned genes and degenerate variants thereof) or polypeptides of the invention (including full length protein, mature protein and truncations or domains thereof), or compounds and other substances that modulate the overall activity of the target gene products, either at the level of target gene/protein expression or target protein activity. Such modulators include polypeptides, analogs, (variants), including fragments and fusion proteins, antibodies and other binding proteins; chemical compounds that directly or indirectly activate or inhibit the polypeptides of the invention (identified, e.g., via drug screening assays as described herein); antisense polynucleotides and polynucleotides suitable for triple helix formation; and in particular antibodies or other binding partners that specifically recognize one or more epitopes of the polypeptides of the invention.

The polypeptides of the present invention may likewise be involved in cellular activation or in one of the other physiological pathways described herein.

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The polynucleotides provided by the present invention can be used by the research community for various purposes. The polynucleotides can be used to express recombinant protein for analysis, characterization or therapeutic use; as markers for tissues in which the corresponding protein is preferentially expressed (either constitutively or at a particular stage of tissue differentiation or development or in disease states); as molecular weight markers on gels; as chromosome markers or tags (when labeled) to identify chromosomes or to map related gene positions: to compare with endogenous DNA sequences in patients to identify potential genetic disorders; as probes to hybridize and thus discover novel, related DNA sequences; as a source of information to derive PCR primers for genetic fingerprinting; as a probe to "subtract-out" known sequences in the process of discovering other novel polynucleotides; for selecting and making oligomers for attachment to a "gene chip" or other support, including for examination of expression patterns; to raise anti-protein antibodies using DNA immunization techniques; and as an antigen to raise anti-DNA antibodies or elicit another immune response. Where the polynucleotide encodes a protein which binds or potentially binds to another protein (such as, for example, in a receptor-ligand interaction), the polynucleotide can also be used in interaction trap assays (such as, for example, that described in Gyuris et al., Cell 75:791-803 (1993)) to identify polynucleotides encoding the other protein with which binding occurs or to identify inhibitors of the binding interaction.

The polypeptides provided by the present invention can similarly be used in assays to determine biological activity, including in a panel of multiple proteins for high-throughput screening; to raise antibodies or to elicit another immune response; as a reagent (including the labeled reagent) in assays designed to quantitatively determine levels of the protein (or its receptor) in biological fluids; as markers for tissues in which the corresponding polypeptide is preferentially expressed (either constitutively or at a particular stage of tissue differentiation or development or in a disease state); and, of course, to isolate correlative receptors or ligands. Proteins involved in these binding interactions can also be used to screen for peptide or small molecule inhibitors or agonists of the binding interaction.

Any or all of these research utilities are capable of being developed into reagent grade or kit format for commercialization as research products.

Methods for performing the uses listed above are well known to those skilled in the art. References disclosing such methods include without limitation "Molecular Cloning: A Laboratory Manual", 2d ed., Cold Spring Harbor Laboratory Press, Sambrook, J., E. F. Fritsch and T. Maniatis eds., 1989, and "Methods in Enzymology: Guide to Molecular Cloning Techniques", Academic Press, Berger, S. L. and A. R. Kimmel eds., 1987.

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4.10.2 NUTRITIONAL USES

Polynucleotides and polypeptides of the present invention can also be used as nutritional sources or supplements. Such uses include without limitation use as a protein or amino acid supplement, use as a carbon source, use as a nitrogen source and use as a source of carbohydrate. In such cases the polypeptide or polynucleotide of the invention can be added to the feed of a particular organism or can be administered as a separate solid or liquid preparation, such as in the form of powder, pills, solutions, suspensions or capsules. In the case of microorganisms, the polypeptide or polynucleotide of the invention can be added to the medium in or on which the microorganism is cultured.

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4.10.3 CYTOKINE AND CELL PROLIFERATION/DIFFERENTIATION ACTIVITY

A polypeptide of the present invention may exhibit activity relating to cytokine, cell proliferation (either inducing or inhibiting) or cell differentiation (either inducing or inhibiting) activity or may induce production of other cytokines in certain cell populations. A polynucleotide of the invention can encode a polypeptide exhibiting such attributes. Many protein factors discovered to date, including all known cytokines, have exhibited activity in one or more factor-dependent cell proliferation assays, and hence the assays serve as a convenient confirmation of cytokine activity. The activity of therapeutic compositions of the present invention is evidenced by any one of a number of routine factor dependent cell proliferation assays for cell lines including, without limitation, 32D, DA2, DA1G, T10, B9, B9/11, BaF3, MC9/G, M+(preB M+), 2E8, RB5, DA1, 123, T1165, HT2, CTLL2, TF-1, Mo7e, CMK, HUVEC, and Caco. Therapeutic compositions of the invention can be used in the following:

Assays for T-cell or thymocyte proliferation include without limitation those described in: Current Protocols in Immunology, Ed by J. E. Coligan, A. M. Kruisbeek, D. H. Margulies, E. M. Shevach, W. Strober, Pub. Greene Publishing Associates and Wiley-Interscience (Chapter 3, *In Vitro* assays for Mouse Lymphocyte Function 3.1-3.19; Chapter 7, Immunologic studies in Humans); Takai et al., J. Immunol. 137:3494-3500, 1986; Bertagnolli et al., J. Immunol. 145:1706-1712, 1990; Bertagnolli et al., Cellular Immunology 133:327-341, 1991; Bertagnolli, et al., I. Immunol. 149:3778-3783, 1992; Bowman et al., I. Immunol. 152:1756-1761, 1994.

Assays for cytokine production and/or proliferation of spleen cells, lymph node cells or thymocytes include, without limitation, those described in: Polyclonal T cell stimulation, Kruisbeek, A. M. and Shevach, E. M. In Current Protocols in Immunology. J. E. e.a. Coligan eds. Vol 1 pp. 3.12.1-3.12.14, John Wiley and Sons, Toronto. 1994; and Measurement of mouse

and human interleukin-y, Schreiber, R. D. In Current Protocols in Immunology. J. E. e.a. Coligan

eds. Vol 1 pp. 6.8.1-6.8.8, John Wiley and Sons, Toronto. 1994.

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Assays for proliferation and differentiation of hematopoietic and lymphopoietic cells include, without limitation, those described in: Measurement of Human and Murine Interleukin 2 and Interleukin 4, Bottomly, K., Davis, L. S. and Lipsky, P. E. In Current Protocols in Immunology. J. E. e.a. Coligan eds. Vol 1 pp. 6.3.1-6.3.12, John Wiley and Sons, Toronto. 1991; deVries et al., J. Exp. Med. 173:1205-1211, 1991; Moreau et al., Nature 336:690-692, 1988; Greenberger et al., Proc. Natl. Acad. Sci. U.S.A. 80:2931-2938, 1983; Measurement of mouse and human interleukin 6--Nordan, R. In Current Protocols in Immunology. J. E. Coligan eds. Vol 1 pp. 6.6.1-6.6.5, John Wiley and Sons, Toronto. 1991; Smith et al., Proc. Natl. Aced. Sci. U.S.A. 83:1857-1861, 1986; Measurement of human Interleukin 11--Bennett, F., Giannotti, J., Clark, S. C. and Turner, K. J. In Current Protocols in Immunology. J. E. Coligan eds. Vol 1 pp. 6.15.1 John Wiley and Sons, Toronto. 1991; Measurement of mouse and human Interleukin 9--Ciarletta, A., Giannotti, J., Clark, S. C. and Turner, K. J. In Current Protocols in Immunology. J. E. Coligan eds. Vol 1 pp. 6.13.1, John Wiley and Sons, Toronto. 1991.

Assays for T-cell clone responses to antigens (which will identify, among others, proteins that affect APC-T cell interactions as well as direct T-cell effects by measuring proliferation and cytokine production) include, without limitation, those described in: Current Protocols in Immunology, Ed by J. E. Coligan, A. M. Kruisbeek, D. H. Margulies, E. M. Shevach, W Strober, Pub. Greene Publishing Associates and Wiley-Interscience (Chapter 3, *In Vitro* assays for Mouse Lymphocyte Function; Chapter 6, Cytokines and their cellular receptors; Chapter 7, Immunologic studies in Humans); Weinberger et al., Proc. Natl. Acad. Sci. USA 77:6091-6095, 1980; Weinberger et al., Eur. J. Immun. 11:405-411, 1981; Takai et al., J. Immunol. 137:3494-3500, 1986; Takai et al., J. Immunol. 140:508-512, 1988.

4.10.4 STEM CELL GROWTH FACTOR ACTIVITY

A polypeptide of the present invention may exhibit stem cell growth factor activity and be involved in the proliferation, differentiation and survival of pluripotent and totipotent stem cells including primordial germ cells, embryonic stem cells, hematopoietic stem cells and/or germ line stem cells. Administration of the polypeptide of the invention to stem cells in vivo or ex vivo is expected to maintain and expand cell populations in a totipotential or pluripotential state which would be useful for re-engineering damaged or diseased tissues, transplantation, manufacture of bio-pharmaceuticals and the development of bio-sensors. The ability to produce large quantities of human cells has important working applications for the production of human proteins which currently must be obtained from non-human sources or donors, implantation of

cells to treat diseases such as Parkinson's, Alzheimer's and other neurodegenerative diseases; tissues for grafting such as bone marrow, skin, cartilage, tendons, bone, muscle (including cardiac muscle), blood vessels, cornea, neural cells, gastrointestinal cells and others; and organs for transplantation such as kidney, liver, pancreas (including islet cells), heart and lung.

It is contemplated that multiple different exogenous growth factors and/or cytokines may be administered in combination with the polypeptide of the invention to achieve the desired effect, including any of the growth factors listed herein, other stem cell maintenance factors, and specifically including stem cell factor (SCF), leukemia inhibitory factor (LIF), Flt-3 ligand (Flt-3L), any of the interleukins, recombinant soluble IL-6 receptor fused to IL-6, macrophage inflammatory protein 1-alpha (MIP-1-alpha), G-CSF, GM-CSF, thrombopoietin (TPO), platelet factor 4 (PF-4), platelet-derived growth factor (PDGF), neural growth factors and basic fibroblast growth factor (bFGF).

Since totipotent stem cells can give rise to virtually any mature cell type, expansion of these cells in culture will facilitate the production of large quantities of mature cells. Techniques for culturing stem cells are known in the art and administration of polypeptides of the invention, optionally with other growth factors and/or cytokines, is expected to enhance the survival and proliferation of the stem cell populations. This can be accomplished by direct administration of the polypeptide of the invention to the culture medium. Alternatively, stroma cells transfected with a polynucleotide that encodes for the polypeptide of the invention can be used as a feeder layer for the stem cell populations in culture or in vivo. Stromal support cells for feeder layers may include embryonic bone marrow fibroblasts, bone marrow stromal cells, fetal liver cells, or cultured embryonic fibroblasts (see U.S. Patent No. 5,690,926).

Stem cells themselves can be transfected with a polynucleotide of the invention to induce autocrine expression of the polypeptide of the invention. This will allow for generation of undifferentiated totipotential/pluripotential stem cell lines that are useful as is or that can then be differentiated into the desired mature cell types. These stable cell lines can also serve as a source of undifferentiated totipotential/pluripotential mRNA to create cDNA libraries and templates for polymerase chain reaction experiments. These studies would allow for the isolation and identification of differentially expressed genes in stem cell populations that regulate stem cell proliferation and/or maintenance.

Expansion and maintenance of totipotent stem cell populations will be useful in the treatment of many pathological conditions. For example, polypeptides of the present invention may be used to manipulate stem cells in culture to give rise to neuroepithelial cells that can be used to augment or replace cells damaged by illness, autoimmune disease, accidental damage or genetic disorders. The polypeptide of the invention may be useful for inducing the proliferation

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of neural cells and for the regeneration of nerve and brain tissue, i.e. for the treatment of central and peripheral nervous system diseases and neuropathies, as well as mechanical and traumatic disorders which involve degeneration, death or trauma to neural cells or nerve tissue. In addition, the expanded stem cell populations can also be genetically altered for gene therapy purposes and to decrease host rejection of replacement tissues after grafting or implantation.

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Expression of the polypeptide of the invention and its effect on stem cells can also be manipulated to achieve controlled differentiation of the stem cells into more differentiated cell types. A broadly applicable method of obtaining pure populations of a specific differentiated cell type from undifferentiated stem cell populations involves the use of a cell-type specific promoter driving a selectable marker. The selectable marker allows only cells of the desired type to survive. For example, stem cells can be induced to differentiate into cardiomyocytes (Wobus et al., Differentiation, 48: 173-182, (1991); Klug et al., J. Clin. Invest., 98(1): 216-224, (1998)) or skeletal muscle cells (Browder, L. W. In: *Principles of Tissue Engineering eds.* Lanza et al., Academic Press (1997)). Alternatively, directed differentiation of stem cells can be accomplished by culturing the stem cells in the presence of a differentiation factor such as retinoic acid and an antagonist of the polypeptide of the invention which would inhibit the effects of endogenous stem cell factor activity and allow differentiation to proceed.

In vitro cultures of stem cells can be used to determine if the polypeptide of the invention exhibits stem cell growth factor activity. Stem cells are isolated from any one of various cell sources (including hematopoietic stem cells and embryonic stem cells) and cultured on a feeder layer, as described by Thompson et al. Proc. Natl. Acad. Sci, U.S.A., 92: 7844-7848 (1995), in the presence of the polypeptide of the invention alone or in combination with other growth factors or cytokines. The ability of the polypeptide of the invention to induce stem cells proliferation is determined by colony formation on semi-solid support e.g. as described by Bernstein et al., Blood, 77: 2316-2321 (1991).

4.10.5 HEMATOPOIESIS REGULATING ACTIVITY

A polypeptide of the present invention may be involved in regulation of hematopoiesis and, consequently, in the treatment of myeloid or lymphoid cell disorders. Even marginal biological activity in support of colony forming cells or of factor-dependent cell lines indicates involvement in regulating hematopoiesis, e.g. in supporting the growth and proliferation of erythroid progenitor cells alone or in combination with other cytokines, thereby indicating utility, for example, in treating various anemias or for use in conjunction with irradiation/chemotherapy to stimulate the production of erythroid precursors and/or erythroid cells; in supporting the growth and proliferation of myeloid cells such as granulocytes and monocytes/macrophages (i.e.,

traditional CSF activity) useful, for example, in conjunction with chemotherapy to prevent or treat consequent myelo-suppression; in supporting the growth and proliferation of megakaryocytes and consequently of platelets thereby allowing prevention or treatment of various platelet disorders such as thrombocytopenia, and generally for use in place of or complimentary to platelet transfusions; and/or in supporting the growth and proliferation of hematopoietic stem cells which are capable of maturing to any and all of the above-mentioned hematopoietic cells and therefore find therapeutic utility in various stem cell disorders (such as those usually treated with transplantation, including, without limitation, aplastic anemia and paroxysmal nocturnal hemoglobinuria), as well as in repopulating the stem cell compartment post irradiation/chemotherapy, either *in-vivo* or *ex-vivo* (i.e., in conjunction with bone marrow transplantation or with peripheral progenitor cell transplantation (homologous or heterologous)) as normal cells or genetically manipulated for gene therapy.

Therapeutic compositions of the invention can be used in the following:

Suitable assays for proliferation and differentiation of various hematopoietic lines are cited above.

Assays for embryonic stem cell differentiation (which will identify, among others, proteins that influence embryonic differentiation hematopoiesis) include, without limitation, those described in: Johansson et al. Cellular Biology 15:141-151, 1995; Keller et al., Molecular and Cellular Biology 13:473-486, 1993; McClanahan et al., Blood 81:2903-2915, 1993.

Assays for stem cell survival and differentiation (which will identify, among others, proteins that regulate lympho-hematopoiesis) include, without limitation, those described in: Methylcellulose colony forming assays, Freshney, M. G. In Culture of Hematopoietic Cells. R. I. Freshney, et al. eds. Vol pp. 265-268, Wiley-Liss, Inc., New York, N.Y. 1994; Hirayama et al., Proc. Natl. Acad. Sci. USA 89:5907-5911, 1992; Primitive hematopoietic colony forming cells with high proliferative potential, McNiece, I. K. and Briddell, R. A. In Culture of Hematopoietic Cells. R. I. Freshney, et al. eds. Vol pp. 23-39, Wiley-Liss, Inc., New York, N.Y. 1994; Neben et al., Experimental Hematology 22:353-359, 1994; Cobblestone area forming cell assay, Ploemacher, R. E. In Culture of Hematopoietic Cells. R. I. Freshney, et al. eds. Vol pp. 1-21, Wiley-Liss, Inc., New York, N.Y. 1994; Long term bone marrow cultures in the presence of stromal cells, Spooncer, E., Dexter, M. and Allen, T. In Culture of Hematopoietic Cells. R. I. Freshney, et al. eds. Vol pp. 163-179, Wiley-Liss, Inc., New York, N.Y. 1994; Long term culture initiating cell assay, Sutherland, H. J. In Culture of Hematopoietic Cells. R. I. Freshney, et al. eds. Vol pp. 139-162, Wiley-Liss, Inc., New York, N.Y. 1994.

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A polypeptide of the present invention also may be involved in bone, cartilage, tendon, ligament and/or nerve tissue growth or regeneration, as well as in wound healing and tissue repair and replacement, and in healing of burns, incisions and ulcers.

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A polypeptide of the present invention which induces cartilage and/or bone growth in circumstances where bone is not normally formed, has application in the healing of bone fractures and cartilage damage or defects in humans and other animals. Compositions of a polypeptide, antibody, binding partner, or other modulator of the invention may have prophylactic use in closed as well as open fracture reduction and also in the improved fixation of artificial joints. De novo bone formation induced by an osteogenic agent contributes to the repair of congenital, trauma induced, or oncologic resection induced craniofacial defects, and also is useful in cosmetic plastic surgery.

A polypeptide of this invention may also be involved in attracting bone-forming cells, stimulating growth of bone-forming cells, or inducing differentiation of progenitors of bone-forming cells. Treatment of osteoporosis, osteoarthritis, bone degenerative disorders, or periodontal disease, such as through stimulation of bone and/or cartilage repair or by blocking inflammation or processes of tissue destruction (collagenase activity, osteoclast activity, etc.) mediated by inflammatory processes may also be possible using the composition of the invention.

Another category of tissue regeneration activity that may involve the polypeptide of the present invention is tendon/ligament formation. Induction of tendon/ligament-like tissue or other tissue formation in circumstances where such tissue is not normally formed, has application in the healing of tendon or ligament tears, deformities and other tendon or ligament defects in humans and other animals. Such a preparation employing a tendon/ligament-like tissue inducing protein may have prophylactic use in preventing damage to tendon or ligament tissue, as well as use in the improved fixation of tendon or ligament to bone or other tissues, and in repairing defects to tendon or ligament tissue. De novo tendon/ligament-like tissue formation induced by a composition of the present invention contributes to the repair of congenital, trauma induced, or other tendon or ligament defects of other origin, and is also useful in cosmetic plastic surgery for attachment or repair of tendons or ligaments. The compositions of the present invention may provide environment to attract tendon- or ligament-forming cells, stimulate growth of tendon- or ligament-forming cells, induce differentiation of progenitors of tendon- or ligament-forming cells, or induce growth of tendon/ligament cells or progenitors ex vivo for return in vivo to effect tissue repair. The compositions of the invention may also be useful in the treatment of tendinitis, carpal tunnel syndrome and other tendon or ligament defects. The compositions may also include an appropriate matrix and/or sequestering agent as a carrier as is well known in the art.

The compositions of the present invention may also be useful for proliferation of neural cells and for regeneration of nerve and brain tissue, i.e. for the treatment of central and peripheral nervous system diseases and neuropathies, as well as mechanical and traumatic disorders, which involve degeneration, death or trauma to neural cells or nerve tissue. More specifically, a composition may be used in the treatment of diseases of the peripheral nervous system, such as peripheral nerve injuries, peripheral neuropathy and localized neuropathies, and central nervous system diseases, such as Alzheimer's, Parkinson's disease, Huntington's disease, amyotrophic lateral sclerosis, and Shy-Drager syndrome. Further conditions which may be treated in accordance with the present invention include mechanical and traumatic disorders, such as spinal cord disorders, head trauma and cerebrovascular diseases such as stroke. Peripheral neuropathies resulting from chemotherapy or other medical therapies may also be treatable using a composition of the invention.

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Compositions of the invention may also be useful to promote better or faster closure of non-healing wounds, including without limitation pressure ulcers, ulcers associated with vascular insufficiency, surgical and traumatic wounds, and the like.

Compositions of the present invention may also be involved in the generation or regeneration of other tissues, such as organs (including, for example, pancreas, liver, intestine, kidney, skin, endothelium), muscle (smooth, skeletal or cardiac) and vascular (including vascular endothelium) tissue, or for promoting the growth of cells comprising such tissues. Part of the desired effects may be by inhibition or modulation of fibrotic scarring may allow normal tissue to regenerate. A polypeptide of the present invention may also exhibit angiogenic activity.

A composition of the present invention may also be useful for gut protection or regeneration and treatment of lung or liver fibrosis, reperfusion injury in various tissues, and conditions resulting from systemic cytokine damage.

A composition of the present invention may also be useful for promoting or inhibiting differentiation of tissues described above from precursor tissues or cells; or for inhibiting the growth of tissues described above.

Therapeutic compositions of the invention can be used in the following:

Assays for tissue generation activity include, without limitation, those described in: International Patent Publication No. WO95/16035 (bone, cartilage, tendon); International Patent Publication No. WO95/05846 (nerve, neuronal); International Patent Publication No. WO91/07491 (skin, endothelium).

Assays for wound healing activity include, without limitation, those described in: Winter, Epidermal Wound Healing, pps. 71-112 (Maibach, H. I. and Rovee, D. T., eds.), Year Book

Medical Publishers, Inc., Chicago, as modified by Eaglstein and Mertz, J. Invest. Dermatol 71:382-84 (1978).

4.10.7 IMMUNE STIMULATING OR SUPPRESSING ACTIVITY

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A polypeptide of the present invention may also exhibit immune stimulating or immune suppressing activity, including without limitation the activities for which assays are described herein. A polynucleotide of the invention can encode a polypeptide exhibiting such activities. A protein may be useful in the treatment of various immune deficiencies and disorders (including severe combined immunodeficiency (SCID)), e.g., in regulating (up or down) growth and proliferation of T and/or B lymphocytes, as well as effecting the cytolytic activity of NK cells and other cell populations. These immune deficiencies may be genetic or be caused by viral (e.g., HIV) as well as bacterial or fungal infections, or may result from autoimmune disorders. More specifically, infectious diseases causes by viral, bacterial, fungal or other infection may be treatable using a protein of the present invention, including infections by HIV, hepatitis viruses, herpes viruses, mycobacteria, Leishmania spp., malaria spp. and various fungal infections such as candidiasis. Of course, in this regard, proteins of the present invention may also be useful where a boost to the immune system generally may be desirable, i.e., in the treatment of cancer.

Autoimmune disorders which may be treated using a protein of the present invention

include, for example, connective tissue disease, multiple sclerosis, systemic lupus erythematosus, rheumatoid arthritis, autoimmune pulmonary inflammation, Guillain-Barre syndrome, autoimmune thyroiditis, insulin dependent diabetes mellitis, myasthenia gravis, graft-versus-host disease and autoimmune inflammatory eye disease. Such a protein (or antagonists thereof, including antibodies) of the present invention may also to be useful in the treatment of allergic reactions and conditions (e.g., anaphylaxis, serum sickness, drug reactions, food allergies, insect venom allergies, mastocytosis, allergic rhinitis, hypersensitivity pneumonitis, urticaria. angioedema, eczema, atopic dermatitis, allergic contact dermatitis, erythema multiforme, Stevens-Johnson syndrome, allergic conjunctivitis, atopic keratoconjunctivitis, venereal keratoconjunctivitis, giant papillary conjunctivitis and contact allergies), such as asthma (particularly allergic asthma) or other respiratory problems. Other conditions, in which immune suppression is desired (including, for example, organ transplantation), may also be treatable using a protein (or antagonists thereof) of the present invention. The therapeutic effects of the polypeptides or antagonists thereof on allergic reactions can be evaluated by in vivo animals models such as the cumulative contact enhancement test (Lastborn et al., Toxicology 125: 59-66, 1998), skin prick test (Hoffmann et al., Allergy 54: 446-54, 1999), guinea pig skin sensitization

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test (Vohr et al., Arch. Toxocol. 73: 501-9), and murine local lymph node assay (Kimber et al., J. Toxicol. Environ. Health 53: 563-79).

Using the proteins of the invention it may also be possible to modulate immune responses, in a number of ways. Down regulation may be in the form of inhibiting or blocking an immune response already in progress or may involve preventing the induction of an immune response. The functions of activated T cells may be inhibited by suppressing T cell responses or by inducing specific tolerance in T cells, or both. Immunosuppression of T cell responses is generally an active, non-antigen-specific, process which requires continuous exposure of the T cells to the suppressive agent. Tolerance, which involves inducing non-responsiveness or anergy in T cells, is distinguishable from immunosuppression in that it is generally antigen-specific and persists after exposure to the tolerizing agent has ceased. Operationally, tolerance can be demonstrated by the lack of a T cell response upon reexposure to specific antigen in the absence of the tolerizing agent.

Down regulating or preventing one or more antigen functions (including without limitation B lymphocyte antigen functions (such as, for example, B7)), e.g., preventing high level lymphokine synthesis by activated T cells, will be useful in situations of tissue, skin and organ transplantation and in graft-versus-host disease (GVHD). For example, blockage of T cell function should result in reduced tissue destruction in tissue transplantation. Typically, in tissue transplants, rejection of the transplant is initiated through its recognition as foreign by T cells, followed by an immune reaction that destroys the transplant. The administration of a therapeutic composition of the invention may prevent cytokine synthesis by immune cells, such as T cells, and thus acts as an immunosuppressant. Moreover, a lack of costimulation may also be sufficient to anergize the T cells, thereby inducing tolerance in a subject. Induction of long-term tolerance by B lymphocyte antigen-blocking reagents may avoid the necessity of repeated administration of these blocking reagents. To achieve sufficient immunosuppression or tolerance in a subject, it may also be necessary to block the function of a combination of B lymphocyte antigens.

The efficacy of particular therapeutic compositions in preventing organ transplant rejection or GVHD can be assessed using animal models that are predictive of efficacy in humans. Examples of appropriate systems which can be used include allogeneic cardiac grafts in rats and xenogeneic pancreatic islet cell grafts in mice, both of which have been used to examine the immunosuppressive effects of CTLA4Ig fusion proteins in vivo as described in Lenschow et al., Science 257:789-792 (1992) and Turka et al., Proc. Natl. Acad. Sci USA, 89:11102-11105 (1992). In addition, murine models of GVHD (see Paul ed., Fundamental Immunology, Raven Press, New York, 1989, pp. 846-847) can be used to determine the effect of therapeutic compositions of the invention on the development of that disease.

Blocking antigen function may also be therapeutically useful for treating autoimmune diseases. Many autoimmune disorders are the result of inappropriate activation of T cells that are reactive against self tissue and which promote the production of cytokines and autoantibodies involved in the pathology of the diseases. Preventing the activation of autoreactive T cells may reduce or eliminate disease symptoms. Administration of reagents which block stimulation of T cells can be used to inhibit T cell activation and prevent production of autoantibodies or T cell-derived cytokines which may be involved in the disease process. Additionally, blocking reagents may induce antigen-specific tolerance of autoreactive T cells which could lead to long-term relief from the disease. The efficacy of blocking reagents in preventing or alleviating autoimmune disorders can be determined using a number of well-characterized animal models of human autoimmune diseases. Examples include murine experimental autoimmune encephalitis, systemic lupus erythmatosis in MRL/lpr/lpr mice or NZB hybrid mice, murine autoimmune collagen arthritis, diabetes mellitus in NOD mice and BB rats, and murine experimental myasthenia gravis (see Paul ed., Fundamental Immunology, Raven Press, New York, 1989, pp. 840-856).

Upregulation of an antigen function (e.g., a B lymphocyte antigen function), as a means of up regulating immune responses, may also be useful in therapy. Upregulation of immune responses may be in the form of enhancing an existing immune response or eliciting an initial immune response. For example, enhancing an immune response may be useful in cases of viral infection, including systemic viral diseases such as influenza, the common cold, and encephalitis.

Alternatively, anti-viral immune responses may be enhanced in an infected patient by removing T cells from the patient, costimulating the T cells in vitro with viral antigen-pulsed APCs either expressing a peptide of the present invention or together with a stimulatory form of a soluble peptide of the present invention and reintroducing the in vitro activated T cells into the patient. Another method of enhancing anti-viral immune responses would be to isolate infected cells from a patient, transfect them with a nucleic acid encoding a protein of the present invention as described herein such that the cells express all or a portion of the protein on their surface, and reintroduce the transfected cells into the patient. The infected cells would now be capable of delivering a costimulatory signal to, and thereby activate, T cells in vivo.

A polypeptide of the present invention may provide the necessary stimulation signal to T cells to induce a T cell mediated immune response against the transfected tumor cells. In addition, tumor cells which lack MHC class I or MHC class II molecules, or which fail to reexpress sufficient mounts of MHC class I or MHC class II molecules, can be transfected with nucleic acid encoding all or a portion of (e.g., a cytoplasmic-domain truncated portion) of an MHC class I alpha chain protein and β₂ microglobulin protein or an MHC class II alpha chain

protein and an MHC class II beta chain protein to thereby express MHC class I or MHC class II proteins on the cell surface. Expression of the appropriate class I or class II MHC in conjunction with a peptide having the activity of a B lymphocyte antigen (e.g., B7-1, B7-2, B7-3) induces a T cell mediated immune response against the transfected tumor cell. Optionally, a gene encoding an antisense construct which blocks expression of an MHC class II associated protein, such as the invariant chain, can also be cotransfected with a DNA encoding a peptide having the activity of a B lymphocyte antigen to promote presentation of tumor associated antigens and induce tumor specific immunity. Thus, the induction of a T cell mediated immune response in a human subject may be sufficient to overcome tumor-specific tolerance in the subject.

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The activity of a protein of the invention may, among other means, be measured by the following methods:

Suitable assays for thymocyte or splenocyte cytotoxicity include, without limitation, those described in: Current Protocols in Immunology, Ed by J. E. Coligan, A. M. Kruisbeek, D. H. Margulies, E. M. Shevach, W. Strober, Pub. Greene Publishing Associates and Wiley-Interscience (Chapter 3, In Vitro assays for Mouse Lymphocyte Function 3.1-3.19; Chapter 7, Immunologic studies in Humans); Herrmann et al., Proc. Natl. Acad. Sci. USA 78:2488-2492, 1981; Herrmann et al., J. Immunol. 128:1968-1974, 1982; Handa et al., J. Immunol. 135:1564-1572, 1985; Takai et al., I. Immunol. 137:3494-3500, 1986; Takai et al., J. Immunol. 140:508-512, 1988; Bowman et al., J. Virology 61:1992-1998; Bertagnolli et al., Cellular Immunology 133:327-341, 1991; Brown et al., J. Immunol. 153:3079-3092, 1994.

Assays for T-cell-dependent immunoglobulin responses and isotype switching (which will identify, among others, proteins that modulate T-cell dependent antibody responses and that affect Th1/Th2 profiles) include, without limitation, those described in: Maliszewski, J. Immunol. 144:3028-3033, 1990; and Assays for B cell function: In vitro antibody production, Mond, J. J. and Brunswick, M. In Current Protocols in Immunology. J. E. e.a. Coligan eds. Vol 1 pp. 3.8.1-3.8.16, John Wiley and Sons, Toronto. 1994.

Mixed lymphocyte reaction (MLR) assays (which will identify, among others, proteins that generate predominantly Th1 and CTL responses) include, without limitation, those described in: Current Protocols in Immunology, Ed by J. E. Coligan, A. M. Kruisbeek, D. H. Margulies, E. M. Shevach, W. Strober, Pub. Greene Publishing Associates and Wiley-Interscience (Chapter 3, In Vitro assays for Mouse Lymphocyte Function 3.1-3.19; Chapter 7, Immunologic studies in Humans); Takai et al., J. Immunol. 137:3494-3500, 1986; Takai et al., J. Immunol. 140:508-512, 1988; Bertagnolli et al., J. Immunol. 149:3778-3783, 1992.

Dendritic cell-dependent assays (which will identify, among others, proteins expressed by dendritic cells that activate naive T-cells) include, without limitation, those described in: Guery

et al., J. Immunol. 134:536-544, 1995; Inaba et al., Journal of Experimental Medicine 173:549-559, 1991; Macatonia et al., Journal of Immunology 154:5071-5079, 1995; Porgador et al., Journal of Experimental Medicine 182:255-260, 1995; Nair et al., Journal of Virology 67:4062-4069, 1993; Huang et al., Science 264:961-965, 1994; Macatonia et al., Journal of Experimental Medicine 169:1255-1264, 1989; Bhardwaj et al., Journal of Clinical Investigation 94:797-807, 1994; and Inaba et al., Journal of Experimental Medicine 172:631-640, 1990.

Assays for lymphocyte survival/apoptosis (which will identify, among others, proteins that prevent apoptosis after superantigen induction and proteins that regulate lymphocyte homeostasis) include, without limitation, those described in: Darzynkiewicz et al., Cytometry 13:795-808, 1992; Gorczyca et al., Leukemia 7:659-670, 1993; Gorczyca et al., Cancer Research 53:1945-1951, 1993; Itoh et al., Cell 66:233-243, 1991; Zacharchuk, Journal of Immunology 145:4037-4045, 1990; Zamai et al., Cytometry 14:891-897, 1993; Gorczyca et al., International Journal of Oncology 1:639-648, 1992.

Assays for proteins that influence early steps of T-cell commitment and development include, without limitation, those described in: Antica et al., Blood 84:111-117, 1994; Fine et al., Cellular Immunology 155:111-122, 1994; Galy et al., Blood 85:2770-2778, 1995; Toki et al., Proc. Nat. Acad Sci. USA 88:7548-7551, 1991.

4.10.8 ACTIVIN/INHIBIN ACTIVITY

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A polypeptide of the present invention may also exhibit activin- or inhibin-related activities. A polynucleotide of the invention may encode a polypeptide exhibiting such characteristics. Inhibins are characterized by their ability to inhibit the release of follicle stimulating hormone (FSH), while activins and are characterized by their ability to stimulate the release of follicle stimulating hormone (FSH). Thus, a polypeptide of the present invention, alone or in heterodimers with a member of the inhibin family, may be useful as a contraceptive based on the ability of inhibins to decrease fertility in female mammals and decrease spermatogenesis in male mammals. Administration of sufficient amounts of other inhibins can induce infertility in these mammals. Alternatively, the polypeptide of the invention, as a homodimer or as a heterodimer with other protein subunits of the inhibin group, may be useful as a fertility inducing therapeutic, based upon the ability of activin molecules in stimulating FSH release from cells of the anterior pituitary. See, for example, U.S. Pat. No. 4,798,885. A polypeptide of the invention may also be useful for advancement of the onset of fertility in sexually immature mammals, so as to increase the lifetime reproductive performance of domestic animals such as, but not limited to, cows, sheep and pigs.

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The activity of a polypeptide of the invention may, among other means, be measured by the following methods.

Assays for activin/inhibin activity include, without limitation, those described in: Vale et al., Endocrinology 91:562-572, 1972; Ling et al., Nature 321:779-782, 1986; Vale et al., Nature 321:776-779, 1986; Mason et al., Nature 318:659-663, 1985; Forage et al., Proc. Natl. Acad. Sci. USA 83:3091-3095, 1986.

4.10.9 CHEMOTACTIC/CHEMOKINETIC ACTIVITY

A polypeptide of the present invention may be involved in chemotactic or chemokinetic activity for mammalian cells, including, for example, monocytes, fibroblasts, neutrophils, T-cells, mast cells, eosinophils, epithelial and/or endothelial cells. A polynucleotide of the invention can encode a polypeptide exhibiting such attributes. Chemotactic and chemokinetic receptor activation can be used to mobilize or attract a desired cell population to a desired site of action. Chemotactic or chemokinetic compositions (e.g. proteins, antibodies, binding partners, or modulators of the invention) provide particular advantages in treatment of wounds and other trauma to tissues, as well as in treatment of localized infections. For example, attraction of lymphocytes, monocytes or neutrophils to tumors or sites of infection may result in improved immune responses against the tumor or infecting agent.

A protein or peptide has chemotactic activity for a particular cell population if it can stimulate, directly or indirectly, the directed orientation or movement of such cell population. Preferably, the protein or peptide has the ability to directly stimulate directed movement of cells. Whether a particular protein has chemotactic activity for a population of cells can be readily determined by employing such protein or peptide in any known assay for cell chemotaxis.

Therapeutic compositions of the invention can be used in the following:

Assays for chemotactic activity (which will identify proteins that induce or prevent chemotaxis) consist of assays that measure the ability of a protein to induce the migration of cells across a membrane as well as the ability of a protein to induce the adhesion of one cell population to another cell population. Suitable assays for movement and adhesion include, without limitation, those described in: Current Protocols in Immunology, Ed by J. E. Coligan, A. M. Kruisbeek, D. H. Marguiles, E. M. Shevach, W. Strober, Pub. Greene Publishing Associates and Wiley-Interscience (Chapter 6.12, Measurement of alpha and beta Chemokines 6.12.1-6.12.28; Taub et al. J. Clin. Invest. 95:1370-1376, 1995; Lind et al. APMIS 103:140-146, 1995; Muller et al Eur. J. Immunol. 25:1744-1748; Gruber et al. J. of Immunol. 152:5860-5867, 1994; Johnston et al. J. of Immunol. 153:1762-1768, 1994.

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4.10.10 HEMOSTATIC AND THROMBOLYTIC ACTIVITY

A polypeptide of the invention may also be involved in hemostatis or thrombolysis or thrombosis. A polynucleotide of the invention can encode a polypeptide exhibiting such attributes. Compositions may be useful in treatment of various coagulation disorders (including hereditary disorders, such as hemophilias) or to enhance coagulation and other hemostatic events in treating wounds resulting from trauma, surgery or other causes. A composition of the invention may also be useful for dissolving or inhibiting formation of thromboses and for treatment and prevention of conditions resulting therefrom (such as, for example, infarction of cardiac and central nervous system vessels (e.g., stroke).

Therapeutic compositions of the invention can be used in the following:

Assay for hemostatic and thrombolytic activity include, without limitation, those described in: Linet et al., J. Clin. Pharmacol. 26:131-140, 1986; Burdick et al., Thrombosis Res. 45:413-419, 1987; Humphrey et al., Fibrinolysis 5:71-79 (1991); Schaub, Prostaglandins 35:467-474, 1988.

4.10.11 CANCER DIAGNOSIS AND THERAPY

Polypeptides of the invention may be involved in cancer cell generation, proliferation or metastasis. Detection of the presence or amount of polynucleotides or polypeptides of the invention may be useful for the diagnosis and/or prognosis of one or more types of cancer. For example, the presence or increased expression of a polynucleotide/polypeptide of the invention may indicate a hereditary risk of cancer, a precancerous condition, or an ongoing malignancy. Conversely, a defect in the gene or absence of the polypeptide may be associated with a cancer condition. Identification of single nucleotide polymorphisms associated with cancer or a predisposition to cancer may also be useful for diagnosis or prognosis.

Cancer treatments promote tumor regression by inhibiting tumor cell proliferation, inhibiting angiogenesis (growth of new blood vessels that is necessary to support tumor growth) and/or prohibiting metastasis by reducing tumor cell motility or invasiveness. Therapeutic compositions of the invention may be effective in adult and pediatric oncology including in solid phase tumors/malignancies, locally advanced tumors, human soft tissue sarcomas, metastatic cancer, including lymphatic metastases, blood cell malignancies including multiple myeloma, acute and chronic leukemias, and lymphomas, head and neck cancers including mouth cancer, larynx cancer and thyroid cancer, lung cancers including small cell carcinoma and non-small cell cancers, breast cancers including small cell carcinoma and ductal carcinoma, gastrointestinal cancers including esophageal cancer, stomach cancer, colon cancer, colorectal cancer and polyps associated with colorectal neoplasia, pancreatic cancers, liver cancer, urologic cancers including

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bladder cancer and prostate cancer, malignancies of the female genital tract including ovarian carcinoma, uterine (including endometrial) cancers, and solid tumor in the ovarian follicle, kidney cancers including renal cell carcinoma, brain cancers including intrinsic brain tumors, neuroblastoma, astrocytic brain tumors, gliomas, metastatic tumor cell invasion in the central nervous system, bone cancers including osteomas, skin cancers including malignant melanoma, tumor progression of human skin keratinocytes, squamous cell carcinoma, basal cell carcinoma, hemangiopericytoma and Karposi's sarcoma.

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Polypeptides, polynucleotides, or modulators of polypeptides of the invention (including inhibitors and stimulators of the biological activity of the polypeptide of the invention) may be administered to treat cancer. Therapeutic compositions can be administered in therapeutically effective dosages alone or in combination with adjuvant cancer therapy such as surgery, chemotherapy, radiotherapy, thermotherapy, and laser therapy, and may provide a beneficial effect, e.g. reducing tumor size, slowing rate of tumor growth, inhibiting metastasis, or otherwise improving overall clinical condition, without necessarily eradicating the cancer.

The composition can also be administered in therapeutically effective amounts as a portion of an anti-cancer cocktail. An anti-cancer cocktail is a mixture of the polypeptide or modulator of the invention with one or more anti-cancer drugs in addition to a pharmaceutically acceptable carrier for delivery. The use of anti-cancer cocktails as a cancer treatment is routine. Anti-cancer drugs that are well known in the art and can be used as a treatment in combination with the polypeptide or modulator of the invention include: Actinomycin D, Aminoglutethimide, Asparaginase, Bleomycin, Busulfan, Carboplatin, Carmustine, Chlorambucil, Cisplatin (cis-DDP), Cyclophosphamide, Cytarabine HCl (Cytosine arabinoside), Dacarbazine, Dactinomycin, Daunorubicin HCl, Doxorubicin HCl, Estramustine phosphate sodium, Etoposide (V16-213), Floxuridine, 5-Fluorouracil (5-Fu), Flutamide, Hydroxyurea (hydroxycarbamide), Ifosfamide, Interferon Alpha-2a, Interferon Alpha-2b, Leuprolide acetate (LHRH-releasing factor analog), Lomustine, Mechlorethamine HCl (nitrogen mustard), Melphalan, Mercaptopurine, Mesna, Methotrexate (MTX), Mitomycin, Mitoxantrone HCl, Octreotide, Plicamycin, Procarbazine HCl, Streptozocin, Tamoxifen citrate, Thioguanine, Thiotepa, Vinblastine sulfate. Vincristine sulfate. Amsacrine, Azacitidine, Hexamethylmelamine, Interleukin-2, Mitoguazone, Pentostatin, Semustine, Teniposide, and Vindesine sulfate.

In addition, therapeutic compositions of the invention may be used for prophylactic treatment of cancer. There are hereditary conditions and/or environmental situations (e.g. exposure to carcinogens) known in the art that predispose an individual to developing cancers. Under these circumstances, it may be beneficial to treat these individuals with therapeutically effective doses of the polypeptide of the invention to reduce the risk of developing cancers.

In vitro models can be used to determine the effective doses of the polypeptide of the invention as a potential cancer treatment. These in vitro models include proliferation assays of cultured tumor cells, growth of cultured tumor cells in soft agar (see Freshney, (1987) Culture of Animal Cells: A Manual of Basic Technique, Wily-Liss, New York, NY Ch 18 and Ch 21), tumor systems in nude mice as described in Giovanella et al., J. Natl. Can. Inst., 52: 921-30 (1974), mobility and invasive potential of tumor cells in Boyden Chamber assays as described in Pilkington et al., Anticancer Res., 17: 4107-9 (1997), and angiogenesis assays such as induction of vascularization of the chick chorioallantoic membrane or induction of vascular endothelial cell migration as described in Ribatta et al., Intl. J. Dev. Biol., 40: 1189-97 (1999) and Li et al.,

Clin. Exp. Metastasis, 17:423-9 (1999), respectively. Suitable tumor cells lines are available.

4.10.12 RECEPTOR/LIGAND ACTIVITY

e.g. from American Type Tissue Culture Collection catalogs.

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A polypeptide of the present invention may also demonstrate activity as receptor, receptor ligand or inhibitor or agonist of receptor/ligand interactions. A polynucleotide of the invention can encode a polypeptide exhibiting such characteristics. Examples of such receptors and ligands include, without limitation, cytokine receptors and their ligands, receptor kinases and their ligands, receptor phosphatases and their ligands, receptors involved in cell-cell interactions and their ligands (including without limitation, cellular adhesion molecules (such as selectins, integrins and their ligands) and receptor/ligand pairs involved in antigen presentation, antigen recognition and development of cellular and humoral immune responses. Receptors and ligands are also useful for screening of potential peptide or small molecule inhibitors of the relevant receptor/ligand interaction. A protein of the present invention (including, without limitation, fragments of receptors and ligands) may themselves be useful as inhibitors of receptor/ligand interactions.

The activity of a polypeptide of the invention may, among other means, be measured by the following methods:

Suitable assays for receptor-ligand activity include without limitation those described in: Current Protocols in Immunology, Ed by J. E. Coligan, A. M. Kruisbeek, D. H. Margulies, E. M. Shevach, W. Strober, Pub. Greene Publishing Associates and Wiley-Interscience (Chapter 7.28, Measurement of Cellular Adhesion under static conditions 7.28.1-7.28.22), Takai et al., Proc. Natl. Acad. Sci. USA 84:6864-6868, 1987; Bierer et al., J. Exp. Med. 168:1145-1156, 1988; Rosenstein et al., J. Exp. Med. 169:149-160 1989; Stoltenborg et al., J. Immunol. Methods 175:59-68, 1994; Stitt et al., Cell 80:661-670, 1995.

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By way of example, the polypeptides of the invention may be used as a receptor for a ligand(s) thereby transmitting the biological activity of that ligand(s). Ligands may be identified through binding assays, affinity chromatography, dihybrid screening assays, BIAcore assays, gel overlay assays, or other methods known in the art.

Studies characterizing drugs or proteins as agonist or antagonist or partial agonists or a partial antagonist require the use of other proteins as competing ligands. The polypeptides of the present invention or ligand(s) thereof may be labeled by being coupled to radioisotopes, colorimetric molecules or a toxin molecules by conventional methods. ("Guide to Protein Purification" Murray P. Deutscher (ed) Methods in Enzymology Vol. 182 (1990) Academic Press, Inc. San Diego). Examples of radioisotopes include, but are not limited to, tritium and carbon-14. Examples of colorimetric molecules include, but are not limited to, fluorescent molecules such as fluorescamine, or rhodamine or other colorimetric molecules. Examples of toxins include, but are not limited, to ricin.

4.10.13 DRUG SCREENING

This invention is particularly useful for screening chemical compounds by using the novel polypeptides or binding fragments thereof in any of a variety of drug screening techniques. The polypeptides or fragments employed in such a test may either be free in solution, affixed to a solid support, borne on a cell surface or located intracellularly. One method of drug screening utilizes eukaryotic or prokaryotic host cells which are stably transformed with recombinant nucleic acids expressing the polypeptide or a fragment thereof. Drugs are screened against such transformed cells in competitive binding assays. Such cells, either in viable or fixed form, can be used for standard binding assays. One may measure, for example, the formation of complexes between polypeptides of the invention or fragments and the agent being tested or examine the diminution in complex formation between the novel polypeptides and an appropriate cell line, which are well known in the art.

Sources for test compounds that may be screened for ability to bind to or modulate (i.e., increase or decrease) the activity of polypeptides of the invention include (1) inorganic and organic chemical libraries, (2) natural product libraries, and (3) combinatorial libraries comprised of either random or mimetic peptides, oligonucleotides or organic molecules.

Chemical libraries may be readily synthesized or purchased from a number of commercial sources, and may include structural analogs of known compounds or compounds that are identified as "hits" or "leads" via natural product screening.

The sources of natural product libraries are microorganisms (including bacteria and fungi), animals, plants or other vegetation, or marine organisms, and libraries of mixtures for

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screening may be created by: (1) fermentation and extraction of broths from soil, plant or marine microorganisms or (2) extraction of the organisms themselves. Natural product libraries include polyketides, non-ribosomal peptides, and (non-naturally occurring) variants thereof. For a review, see *Science 282*:63-68 (1998).

Combinatorial libraries are composed of large numbers of peptides, oligonucleotides or organic compounds and can be readily prepared by traditional automated synthesis methods, PCR, cloning or proprietary synthetic methods. Of particular interest are peptide and oligonucleotide combinatorial libraries. Still other libraries of interest include peptide, protein, peptidomimetic, multiparallel synthetic collection, recombinatorial, and polypeptide libraries. For a review of combinatorial chemistry and libraries created therefrom, see Myers, *Curr. Opin. Biotechnol.* 8:701-707 (1997). For reviews and examples of peptidomimetic libraries, see Al-Obeidi et al., *Mol. Biotechnol*, 9(3):205-23 (1998); Hruby et al., *Curr Opin Chem Biol*, 1(1):114-19 (1997); Dorner et al., *Bioorg Med Chem*, 4(5):709-15 (1996) (alkylated dipeptides).

Identification of modulators through use of the various libraries described herein permits modification of the candidate "hit" (or "lead") to optimize the capacity of the "hit" to bind a polypeptide of the invention. The molecules identified in the binding assay are then tested for antagonist or agonist activity in *in vivo* tissue culture or animal models that are well known in the art. In brief, the molecules are titrated into a plurality of cell cultures or animals and then tested for either cell/animal death or prolonged survival of the animal/cells.

The binding molecules thus identified may be complexed with toxins, e.g., ricin or cholera, or with other compounds that are toxic to cells such as radioisotopes. The toxin-binding molecule complex is then targeted to a tumor or other cell by the specificity of the binding molecule for a polypeptide of the invention. Alternatively, the binding molecules may be complexed with imaging agents for targeting and imaging purposes.

4.10.14 ASSAY FOR RECEPTOR ACTIVITY

The invention also provides methods to detect specific binding of a polypeptide e.g. a ligand or a receptor. The art provides numerous assays particularly useful for identifying previously unknown binding partners for receptor polypeptides of the invention. For example, expression cloning using mammalian or bacterial cells, or dihybrid screening assays can be used to identify polynucleotides encoding binding partners. As another example, affinity chromatography with the appropriate immobilized polypeptide of the invention can be used to isolate polypeptides that recognize and bind polypeptides of the invention. There are a number of different libraries used for the identification of compounds, and in particular small molecules, that modulate (*i.e.*, increase or decrease) biological activity of a polypeptide of the invention.

Ligands for receptor polypeptides of the invention can also be identified by adding exogenous ligands, or cocktails of ligands to two cells populations that are genetically identical except for the expression of the receptor of the invention: one cell population expresses the receptor of the invention whereas the other does not. The response of the two cell populations to the addition of ligands(s) are then compared. Alternatively, an expression library can be co-expressed with the polypeptide of the invention in cells and assayed for an autocrine response to identify potential ligand(s). As still another example, BIAcore assays, gel overlay assays, or other methods known in the art can be used to identify binding partner polypeptides, including, (1) organic and inorganic chemical libraries, (2) natural product libraries, and (3) combinatorial libraries comprised of random peptides, oligonucleotides or organic molecules.

The role of downstream intracellular signaling molecules in the signaling cascade of the polypeptide of the invention can be determined. For example, a chimeric protein in which the cytoplasmic domain of the polypeptide of the invention is fused to the extracellular portion of a protein, whose ligand has been identified, is produced in a host cell. The cell is then incubated with the ligand specific for the extracellular portion of the chimeric protein, thereby activating the chimeric receptor. Known downstream proteins involved in intracellular signaling can then be assayed for expected modifications i.e. phosphorylation. Other methods known to those in the art can also be used to identify signaling molecules involved in receptor activity.

4.10.15 ANTI-INFLAMMATORY ACTIVITY

Compositions of the present invention may also exhibit anti-inflammatory activity. The anti-inflammatory activity may be achieved by providing a stimulus to cells involved in the inflammatory response, by inhibiting or promoting cell-cell interactions (such as, for example, cell adhesion), by inhibiting or promoting chemotaxis of cells involved in the inflammatory process, inhibiting or promoting cell extravasation, or by stimulating or suppressing production of other factors which more directly inhibit or promote an inflammatory response. Compositions with such activities can be used to treat inflammatory conditions including chronic or acute conditions), including without limitation intimation associated with infection (such as septic shock, sepsis or systemic inflammatory response syndrome (SIRS)), ischemia-reperfusion injury, endotoxin lethality, arthritis, complement-mediated hyperacute rejection, nephritis, cytokine or chemokine-induced lung injury, inflammatory bowel disease, Crohn's disease or resulting from over production of cytokines such as TNF or IL-1. Compositions of the invention may also be useful to treat anaphylaxis and hypersensitivity to an antigenic substance or material.

Compositions of this invention may be utilized to prevent or treat conditions such as, but not limited to, sepsis, acute pancreatitis, endotoxin shock, cytokine induced shock, rheumatoid

arthritis, chronic inflammatory arthritis, pancreatic cell damage from diabetes mellitus type 1, graft versus host disease, inflammatory bowel disease, inflamation associated with pulmonary disease, other autoimmune disease or inflammatory disease, an antiproliferative agent such as for acute or chronic mylegenous leukemia or in the prevention of premature labor secondary to

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intrauterine infections.

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Leukemias and related disorders may be treated or prevented by administration of a therapeutic that promotes or inhibits function of the polynucleotides and/or polypeptides of the invention. Such leukemias and related disorders include but are not limited to acute leukemia, acute lymphocytic leukemia, acute myelocytic leukemia, myeloblastic, promyelocytic, myelomonocytic, monocytic, erythroleukemia, chronic leukemia, chronic myelocytic (granulocytic) leukemia and chronic lymphocytic leukemia (for a review of such disorders, see Fishman et al., 1985, Medicine, 2d Ed., J.B. Lippincott Co., Philadelphia).

4.10.17 NERVOUS SYSTEM DISORDERS

Nervous system disorders, involving cell types which can be tested for efficacy of intervention with compounds that modulate the activity of the polynucleotides and/or polypeptides of the invention, and which can be treated upon thus observing an indication of therapeutic utility, include but are not limited to nervous system injuries, and diseases or disorders which result in either a disconnection of axons, a diminution or degeneration of neurons, or demyelination. Nervous system lesions which may be treated in a patient (including human and non-human mammalian patients) according to the invention include but are not limited to the following lesions of either the central (including spinal cord, brain) or peripheral nervous systems:

- (i) traumatic lesions, including lesions caused by physical injury or associated with surgery, for example, lesions which sever a portion of the nervous system, or compression injuries;
- (ii) ischemic lesions, in which a lack of oxygen in a portion of the nervous system results in neuronal injury or death, including cerebral infarction or ischemia, or spinal cord infarction or ischemia;
 - (iii) infectious lesions, in which a portion of the nervous system is destroyed or injured as a result of infection, for example, by an abscess or associated with infection by human immunodeficiency virus, herpes zoster, or herpes simplex virus or with Lyme disease, tuberculosis, syphilis;

(iv) degenerative lesions, in which a portion of the nervous system is destroyed or injured as a result of a degenerative process including but not limited to degeneration associated with Parkinson's disease, Alzheimer's disease, Huntington's chorea, or amyotrophic lateral sclerosis;

(v) lesions associated with nutritional diseases or disorders, in which a portion of the nervous system is destroyed or injured by a nutritional disorder or disorder of metabolism including but not limited to, vitamin B12 deficiency, folic acid deficiency, Wernicke disease, tobacco-alcohol amblyopia, Marchiafava-Bignami disease (primary degeneration of the corpus callosum), and alcoholic cerebellar degeneration;

- (vi) neurological lesions associated with systemic diseases including but not limited to diabetes (diabetic neuropathy, Bell's palsy), systemic lupus erythematosus, carcinoma, or sarcoidosis;
- (vii) lesions caused by toxic substances including alcohol, lead, or particular neurotoxins; and
- (viii) demyelinated lesions in which a portion of the nervous system is destroyed or injured by a demyelinating disease including but not limited to multiple sclerosis, human immunodeficiency virus-associated myelopathy, transverse myelopathy or various etiologies, progressive multifocal leukoencephalopathy, and central pontine myelinolysis.

Therapeutics which are useful according to the invention for treatment of a nervous system disorder may be selected by testing for biological activity in promoting the survival or differentiation of neurons. For example, and not by way of limitation, therapeutics which elicit any of the following effects may be useful according to the invention:

(i) increased survival time of neurons in culture;

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- (ii) increased sprouting of neurons in culture or in vivo;
- (iii) increased production of a neuron-associated molecule in culture or *in vivo*, *e.g.*, choline acetyltransferase or acetylcholinesterase with respect to motor neurons; or
 - (iv) decreased symptoms of neuron dysfunction in vivo.

Such effects may be measured by any method known in the art. In preferred, non-limiting embodiments, increased survival of neurons may be measured by the method set forth in Arakawa et al. (1990, J. Neurosci. 10:3507-3515); increased sprouting of neurons may be detected by methods set forth in Pestronk et al. (1980, Exp. Neurol. 70:65-82) or Brown et al. (1981, Ann. Rev. Neurosci. 4:17-42); increased production of neuron-associated molecules may be measured by bioassay, enzymatic assay, antibody binding, Northern blot assay, etc., depending on the molecule to be measured; and motor neuron dysfunction may be measured by

assessing the physical manifestation of motor neuron disorder, e.g., weakness, motor neuron conduction velocity, or functional disability.

In specific embodiments, motor neuron disorders that may be treated according to the invention include but are not limited to disorders such as infarction, infection, exposure to toxin, trauma, surgical damage, degenerative disease or malignancy that may affect motor neurons as well as other components of the nervous system, as well as disorders that selectively affect neurons such as amyotrophic lateral sclerosis, and including but not limited to progressive spinal muscular atrophy, progressive bulbar palsy, primary lateral sclerosis, infantile and juvenile muscular atrophy, progressive bulbar paralysis of childhood (Fazio-Londe syndrome), poliomyelitis and the post polio syndrome, and Hereditary Motorsensory Neuropathy (Charcot-Marie-Tooth Disease).

4.10.18 OTHER ACTIVITIES

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A polypeptide of the invention may also exhibit one or more of the following additional activities or effects: inhibiting the growth, infection or function of, or killing, infectious agents. including, without limitation, bacteria, viruses, fungi and other parasites; effecting (suppressing or enhancing) bodily characteristics, including, without limitation, height, weight, hair color, eye color, skin, fat to lean ratio or other tissue pigmentation, or organ or body part size or shape (such as, for example, breast augmentation or diminution, change in bone form or shape); effecting biorhythms or circadian cycles or rhythms; effecting the fertility of male or female subjects; effecting the metabolism, catabolism, anabolism, processing, utilization, storage or elimination of dietary fat, lipid, protein, carbohydrate, vitamins, minerals, co-factors or other nutritional factors or component(s); effecting behavioral characteristics, including, without limitation, appetite, libido, stress, cognition (including cognitive disorders), depression (including depressive disorders) and violent behaviors; providing analgesic effects or other pain reducing effects; promoting differentiation and growth of embryonic stem cells in lineages other than hematopoietic lineages; hormonal or endocrine activity; in the case of enzymes, correcting deficiencies of the enzyme and treating deficiency-related diseases; treatment of hyperproliferative disorders (such as, for example, psoriasis); immunoglobulin-like activity (such as, for example, the ability to bind antigens or complement); and the ability to act as an antigen in a vaccine composition to raise an immune response against such protein or another material or entity which is cross-reactive with such protein.

4.10.19 IDENTIFICATION OF POLYMORPHISMS

The demonstration of polymorphisms makes possible the identification of such polymorphisms in human subjects and the pharmacogenetic use of this information for diagnosis and treatment. Such polymorphisms may be associated with, e.g., differential predisposition or susceptibility to various disease states (such as disorders involving inflammation or immune response) or a differential response to drug administration, and this genetic information can be used to tailor preventive or therapeutic treatment appropriately. For example, the existence of a polymorphism associated with a predisposition to inflammation or autoimmune disease makes possible the diagnosis of this condition in humans by identifying the presence of the polymorphism.

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Polymorphisms can be identified in a variety of ways known in the art which all generally involve obtaining a sample from a patient, analyzing DNA from the sample, optionally involving isolation or amplification of the DNA, and identifying the presence of the polymorphism in the DNA. For example, PCR may be used to amplify an appropriate fragment of genomic DNA which may then be sequenced. Alternatively, the DNA may be subjected to allele-specific oligonucleotide hybridization (in which appropriate oligonucleotides are hybridized to the DNA under conditions permitting detection of a single base mismatch) or to a single nucleotide extension assay (in which an oligonucleotide that hybridizes immediately adjacent to the position of the polymorphism is extended with one or more labeled nucleotides). In addition, traditional restriction fragment length polymorphism analysis (using restriction enzymes that provide differential digestion of the genomic DNA depending on the presence or absence of the polymorphism) may be performed. Arrays with nucleotide sequences of the present invention can be used to detect polymorphisms. The array can comprise modified nucleotide sequences of the present invention in order to detect the nucleotide sequences of the present invention. In the alternative, any one of the nucleotide sequences of the present invention can be placed on the array to detect changes from those sequences.

Alternatively a polymorphism resulting in a change in the amino acid sequence could also be detected by detecting a corresponding change in amino acid sequence of the protein, e.g., by an antibody specific to the variant sequence.

4.10.20 ARTHRITIS AND INFLAMMATION

The immunosuppressive effects of the compositions of the invention against rheumatoid arthritis is determined in an experimental animal model system. The experimental model system is adjuvant induced arthritis in rats, and the protocol is described by J. Holoshitz, et at., 1983, Science, 219:56, or by B. Waksman et al., 1963, Int. Arch. Allergy Appl. Immunol., 23:129. Induction of the disease can be caused by a single injection, generally intradermally, of a

suspension of killed Mycobacterium tuberculosis in complete Freund's adjuvant (CFA). The route of injection can vary, but rats may be injected at the base of the tail with an adjuvant mixture. The polypeptide is administered in phosphate buffered solution (PBS) at a dose of about 1-5 mg/kg. The control consists of administering PBS only.

The procedure for testing the effects of the test compound would consist of intradermally injecting killed Mycobacterium tuberculosis in CFA followed by immediately administering the test compound and subsequent treatment every other day until day 24. At 14, 15, 18, 20, 22, and 24 days after injection of Mycobacterium CFA, an overall arthritis score may be obtained as described by J. Holoskitz above. An analysis of the data would reveal that the test compound would have a dramatic affect on the swelling of the joints as measured by a decrease of the arthritis score.

4.11 THERAPEUTIC METHODS

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The compositions (including polypeptide fragments, analogs, variants and antibodies or other binding partners or modulators including antisense polynucleotides) of the invention have numerous applications in a variety of therapeutic methods. Examples of therapeutic applications include, but are not limited to, those exemplified herein.

4.11.1 EXAMPLE

One embodiment of the invention is the administration of an effective amount of the polypeptides or other composition of the invention to individuals affected by a disease or disorder that can be modulated by regulating the peptides of the invention. While the mode of administration is not particularly important, parenteral administration is preferred. An exemplary mode of administration is to deliver an intravenous bolus. The dosage of the polypeptides or other composition of the invention will normally be determined by the prescribing physician. It is to be expected that the dosage will vary according to the age, weight, condition and response of the individual patient. Typically, the amount of polypeptide administered per dose will be in the range of about 0.01µg/kg to 100 mg/kg of body weight, with the preferred dose being about 0.1µg/kg to 10 mg/kg of patient body weight. For parenteral administration, polypeptides of the invention will be formulated in an injectable form combined with a pharmaceutically acceptable parenteral vehicle. Such vehicles are well known in the art and examples include water, saline, Ringer's solution, dextrose solution, and solutions consisting of small amounts of the human serum albumin. The vehicle may contain minor amounts of additives that maintain the isotonicity and stability of the polypeptide or other active ingredient. The preparation of such solutions is within the skill of the art.

4.12 PHARMACEUTICAL FORMULATIONS AND ROUTES OF ADMINISTRATION

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A protein or other composition of the present invention (from whatever source derived, including without limitation from recombinant and non-recombinant sources and including antibodies and other binding partners of the polypeptides of the invention) may be administered to a patient in need, by itself, or in pharmaceutical compositions where it is mixed with suitable carriers or excipient(s) at doses to treat or ameliorate a variety of disorders. Such a composition may optionally contain (in addition to protein or other active ingredient and a carrier) diluents, fillers, salts, buffers, stabilizers, solubilizers, and other materials well known in the art. The term "pharmaceutically acceptable" means a non-toxic material that does not interfere with the effectiveness of the biological activity of the active ingredient(s). The characteristics of the carrier will depend on the route of administration. The pharmaceutical composition of the invention may also contain cytokines, lymphokines, or other hematopoietic factors such as M-CSF, GM-CSF, TNF, IL-1, IL-2, IL-3, IL-4, IL-5, IL-6, IL-7, IL-8, IL-9, IL-10, IL-11, IL-12, IL-13, IL-14, IL-15, IFN, TNF0, TNF1, TNF2, G-CSF, Meg-CSF, thrombopoietin, stem cell factor, and erythropoietin. In further compositions, proteins of the invention may be combined with other agents beneficial to the treatment of the disease or disorder in question. These agents include various growth factors such as epidermal growth factor (EGF), platelet-derived growth factor (PDGF), transforming growth factors (TGF-α and TGF-β), insulin-like growth factor (IGF), as well as cytokines described herein.

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The pharmaceutical composition may further contain other agents which either enhance the activity of the protein or other active ingredient or complement its activity or use in treatment. Such additional factors and/or agents may be included in the pharmaceutical composition to produce a synergistic effect with protein or other active ingredient of the invention, or to minimize side effects. Conversely, protein or other active ingredient of the present invention may be included in formulations of the particular clotting factor, cytokine, lymphokine, other hematopoietic factor, thrombolytic or anti-thrombotic factor, or anti-inflammatory agent to minimize side effects of the clotting factor, cytokine, lymphokine, other hematopoietic factor, thrombolytic or anti-thrombotic factor, or anti-inflammatory agent (such as IL-1Ra, IL-1 Hy1, IL-1 Hy2, anti-TNF, corticosteroids, immunosuppressive agents). A protein of the present invention may be active in multimers (e.g., heterodimers or homodimers) or complexes with itself or other proteins. As a result, pharmaceutical compositions of the invention may comprise a protein of the invention in such multimeric or complexed form.

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As an alternative to being included in a pharmaceutical composition of the invention including a first protein, a second protein or a therapeutic agent may be concurrently administered with the first protein (e.g., at the same time, or at differing times provided that therapeutic concentrations of the combination of agents is achieved at the treatment site). Techniques for formulation and administration of the compounds of the instant application may be found in "Remington's Pharmaceutical Sciences," Mack Publishing Co., Easton, PA, latest edition. A therapeutically effective dose further refers to that amount of the compound sufficient to result in amelioration of symptoms, e.g., treatment, healing, prevention or amelioration of the relevant medical condition, or an increase in rate of treatment, healing, prevention or amelioration of such conditions. When applied to an individual active ingredient, administered alone, a therapeutically effective dose refers to that ingredient alone. When applied to a combination, a therapeutically effective dose refers to combined amounts of the active ingredients that result in the therapeutic effect, whether administered in combination, serially or simultaneously.

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In practicing the method of treatment or use of the present invention, a therapeutically effective amount of protein or other active ingredient of the present invention is administered to a mammal having a condition to be treated. Protein or other active ingredient of the present invention may be administered in accordance with the method of the invention either alone or in combination with other therapies such as treatments employing cytokines, lymphokines or other hematopoietic factors. When co- administered with one or more cytokines, lymphokines or other hematopoietic factors, protein or other active ingredient of the present invention may be administered either simultaneously with the cytokine(s), lymphokine(s), other hematopoietic factor(s), thrombolytic or anti-thrombotic factors, or sequentially. If administered sequentially, the attending physician will decide on the appropriate sequence of administering protein or other active ingredient of the present invention in combination with cytokine(s), lymphokine(s), other hematopoietic factor(s), thrombolytic or anti-thrombotic factors.

4.12.1 ROUTES OF ADMINISTRATION

Suitable routes of administration may, for example, include oral, rectal, transmucosal, or intestinal administration; parenteral delivery, including intramuscular, subcutaneous, intramedullary injections, as well as intrathecal, direct intraventricular, intravenous, intraperitoneal, intranasal, or intraocular injections. Administration of protein or other active ingredient of the present invention used in the pharmaceutical composition or to practice the method of the present invention can be carried out in a variety of conventional ways, such as oral

ingestion, inhalation, topical application or cutaneous, subcutaneous, intraperitoneal, parenteral

or intravenous injection. Intravenous administration to the patient is preferred.

Alternately, one may administer the compound in a local rather than systemic manner, for example, via injection of the compound directly into a arthritic joints or in fibrotic tissue, often in a depot or sustained release formulation. In order to prevent the scarring process frequently occurring as complication of glaucoma surgery, the compounds may be administered topically, for example, as eye drops. Furthermore, one may administer the drug in a targeted drug delivery system, for example, in a liposome coated with a specific antibody, targeting, for example, arthritic or fibrotic tissue. The liposomes will be targeted to and taken up selectively by the

The polypeptides of the invention are administered by any route that delivers an effective dosage to the desired site of action. The determination of a suitable route of administration and an effective dosage for a particular indication is within the level of skill in the art. Preferably for wound treatment, one administers the therapeutic compound directly to the site. Suitable dosage ranges for the polypeptides of the invention can be extrapolated from these dosages or from similar studies in appropriate animal models. Dosages can then be adjusted as necessary by the clinician to provide maximal therapeutic benefit.

4.12.2 COMPOSITIONS/FORMULATIONS

Pharmaceutical compositions for use in accordance with the present invention thus may be formulated in a conventional manner using one or more physiologically acceptable carriers comprising excipients and auxiliaries which facilitate processing of the active compounds into preparations which can be used pharmaceutically. These pharmaceutical compositions may be manufactured in a manner that is itself known, e.g., by means of conventional mixing, dissolving, granulating, dragee-making, levigating, emulsifying, encapsulating, entrapping or lyophilizing processes. Proper formulation is dependent upon the route of administration chosen. When a therapeutically effective amount of protein or other active ingredient of the present invention is administered orally, protein or other active ingredient of the present invention will be in the form of a tablet, capsule, powder, solution or elixir. When administered in tablet form, the pharmaceutical composition of the invention may additionally contain a solid carrier such as a gelatin or an adjuvant. The tablet, capsule, and powder contain from about 5 to 95% protein or other active ingredient of the present invention, and preferably from about 25 to 90% protein or other active ingredient of the present invention. When administered in liquid form, a liquid carrier such as water, petroleum, oils of animal or plant origin such as peanut oil, mineral oil, soybean oil, or sesame oil, or synthetic oils may be added. The liquid form of the

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afflicted tissue.

pharmaceutical composition may further contain physiological saline solution, dextrose or other saccharide solution, or glycols such as ethylene glycol, propylene glycol or polyethylene glycol. When administered in liquid form, the pharmaceutical composition contains from about 0.5 to 90% by weight of protein or other active ingredient of the present invention, and preferably from about 1 to 50% protein or other active ingredient of the present invention.

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When a therapeutically effective amount of protein or other active ingredient of the present invention is administered by intravenous, cutaneous or subcutaneous injection, protein or other active ingredient of the present invention will be in the form of a pyrogen-free, parenterally acceptable aqueous solution. The preparation of such parenterally acceptable protein or other active ingredient solutions, having due regard to pH, isotonicity, stability, and the like, is within the skill in the art. A preferred pharmaceutical composition for intravenous, cutaneous, or subcutaneous injection should contain, in addition to protein or other active ingredient of the present invention, an isotonic vehicle such as Sodium Chloride Injection, Ringer's Injection, Dextrose Injection, Dextrose and Sodium Chloride Injection, Lactated Ringer's Injection, or other vehicle as known in the art. The pharmaceutical composition of the present invention may also contain stabilizers, preservatives, buffers, antioxidants, or other additives known to those of skill in the art. For injection, the agents of the invention may be formulated in aqueous solutions, preferably in physiologically compatible buffers such as Hanks's solution, Ringer's solution, or physiological saline buffer. For transmucosal administration, penetrants appropriate to the barrier to be permeated are used in the formulation. Such penetrants are generally known in the art.

For oral administration, the compounds can be formulated readily by combining the active compounds with pharmaceutically acceptable carriers well known in the art. Such carriers enable the compounds of the invention to be formulated as tablets, pills, dragees, capsules, liquids, gels, syrups, slurries, suspensions and the like, for oral ingestion by a patient to be treated. Pharmaceutical preparations for oral use can be obtained from a solid excipient, optionally grinding a resulting mixture, and processing the mixture of granules, after adding suitable auxiliaries, if desired, to obtain tablets or dragee cores. Suitable excipients are, in particular, fillers such as sugars, including lactose, sucrose, mannitol, or sorbitol; cellulose preparations such as, for example, maize starch, wheat starch, rice starch, potato starch, gelatin, gum tragacanth, methyl cellulose, hydroxypropylmethyl-cellulose, sodium carboxymethylcellulose, and/or polyvinylpyrrolidone (PVP). If desired, disintegrating agents may be added, such as the cross-linked polyvinyl pyrrolidone, agar, or alginic acid or a salt thereof such as sodium alginate. Dragee cores are provided with suitable coatings. For this purpose, concentrated sugar solutions may be used, which may optionally contain gum arabic,

talc, polyvinyl pyrrolidone, carbopol gel, polyethylene glycol, and/or titanium dioxide, lacquer solutions, and suitable organic solvents or solvent mixtures. Dyestuffs or pigments may be added to the tablets or dragee coatings for identification or to characterize different combinations of active compound doses.

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Pharmaceutical preparations which can be used orally include push-fit capsules made of gelatin, as well as soft, sealed capsules made of gelatin and a plasticizer, such as glycerol or sorbitol. The push-fit capsules can contain the active ingredients in admixture with filler such as lactose, binders such as starches, and/or lubricants such as talc or magnesium stearate and, optionally, stabilizers. In soft capsules, the active compounds may be dissolved or suspended in suitable liquids, such as fatty oils, liquid paraffin, or liquid polyethylene glycols. In addition, stabilizers may be added. All formulations for oral administration should be in dosages suitable for such administration. For buccal administration, the compositions may take the form of tablets or lozenges formulated in conventional manner.

For administration by inhalation, the compounds for use according to the present invention are conveniently delivered in the form of an aerosol spray presentation from pressurized packs or a nebuliser, with the use of a suitable propellant, e.g., dichlorodifluoromethane, trichlorofluoromethane, dichlorotetrafluoroethane, carbon dioxide or other suitable gas. In the case of a pressurized aerosol the dosage unit may be determined by providing a valve to deliver a metered amount. Capsules and cartridges of, e.g., gelatin for use in an inhaler or insufflator may be formulated containing a powder mix of the compound and a suitable powder base such as lactose or starch. The compounds may be formulated for parenteral administration by injection, e.g., by bolus injection or continuous infusion. Formulations for injection may be presented in unit dosage form, e.g., in ampules or in multi-dose containers, with an added preservative. The compositions may take such forms as suspensions, solutions or emulsions in oily or aqueous vehicles, and may contain formulatory agents such as suspending, stabilizing and/or dispersing agents.

Pharmaceutical formulations for parenteral administration include aqueous solutions of the active compounds in water-soluble form. Additionally, suspensions of the active compounds may be prepared as appropriate oily injection suspensions. Suitable lipophilic solvents or vehicles include fatty oils such as sesame oil, or synthetic fatty acid esters, such as ethyl oleate or triglycerides, or liposomes. Aqueous injection suspensions may contain substances which increase the viscosity of the suspension, such as sodium carboxymethyl cellulose, sorbitol, or dextran. Optionally, the suspension may also contain suitable stabilizers or agents which increase the solubility of the compounds to allow for the preparation of highly concentrated

solutions. Alternatively, the active ingredient may be in powder form for constitution with a

suitable vehicle, e.g., sterile pyrogen-free water, before use.

sparingly soluble derivatives, for example, as a sparingly soluble salt.

The compounds may also be formulated in rectal compositions such as suppositories or retention enemas, e.g., containing conventional suppository bases such as cocoa butter or other glycerides. In addition to the formulations described previously, the compounds may also be formulated as a depot preparation. Such long acting formulations may be administered by implantation (for example subcutaneously or intramuscularly) or by intramuscular injection. Thus, for example, the compounds may be formulated with suitable polymeric or hydrophobic materials (for example as an emulsion in an acceptable oil) or ion exchange resins, or as

A pharmaceutical carrier for the hydrophobic compounds of the invention is a co-solvent system comprising benzyl alcohol, a nonpolar surfactant, a water-miscible organic polymer, and an aqueous phase. The co-solvent system may be the VPD co-solvent system. VPD is a solution of 3% w/v benzyl alcohol, 8% w/v of the nonpolar surfactant polysorbate 80, and 65% w/v polyethylene glycol 300, made up to volume in absolute ethanol. The VPD co-solvent system (VPD:5W) consists of VPD diluted 1:1 with a 5% dextrose in water solution. This co-solvent system dissolves hydrophobic compounds well, and itself produces low toxicity upon systemic administration. Naturally, the proportions of a co-solvent system may be varied considerably without destroying its solubility and toxicity characteristics. Furthermore, the identity of the co-solvent components may be varied: for example, other low-toxicity nonpolar surfactants may be used instead of polysorbate 80; the fraction size of polyethylene glycol may be varied; other biocompatible polymers may replace polyethylene glycol, e.g. polyvinyl pyrrolidone; and other sugars or polysaccharides may substitute for dextrose. Alternatively, other delivery systems for hydrophobic pharmaceutical compounds may be employed. Liposomes and emulsions are well known examples of delivery vehicles or carriers for hydrophobic drugs. Certain organic solvents such as dimethylsulfoxide also may be employed, although usually at the cost of greater toxicity. Additionally, the compounds may be delivered using a sustained-release system, such as semipermeable matrices of solid hydrophobic polymers containing the therapeutic agent. Various types of sustained-release materials have been established and are well known by those skilled in the art. Sustained-release capsules may, depending on their chemical nature, release the compounds for a few weeks up to over 100 days. Depending on the chemical nature and the biological stability of the therapeutic reagent, additional strategies for protein or other active ingredient stabilization may be employed.

The pharmaceutical compositions also may comprise suitable solid or gel phase carriers or excipients. Examples of such carriers or excipients include but are not limited to calcium

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carbonate, calcium phosphate, various sugars, starches, cellulose derivatives, gelatin, and polymers such as polyethylene glycols. Many of the active ingredients of the invention may be provided as salts with pharmaceutically compatible counter ions. Such pharmaceutically acceptable base addition salts are those salts which retain the biological effectiveness and properties of the free acids and which are obtained by reaction with inorganic or organic bases such as sodium hydroxide, magnesium hydroxide, ammonia, trialkylamine, dialkylamine, monoalkylamine, dibasic amino acids, sodium acetate, potassium benzoate, triethanol amine and the like.

The pharmaceutical composition of the invention may be in the form of a complex of the protein(s) or other active ingredient(s) of present invention along with protein or peptide antigens. The protein and/or peptide antigen will deliver a stimulatory signal to both B and T lymphocytes. B lymphocytes will respond to antigen through their surface immunoglobulin receptor. T lymphocytes will respond to antigen through the T cell receptor (TCR) following presentation of the antigen by MHC proteins. MHC and structurally related proteins including those encoded by class I and class II MHC genes on host cells will serve to present the peptide antigen(s) to T lymphocytes. The antigen components could also be supplied as purified MHC-peptide complexes alone or with co-stimulatory molecules that can directly signal T cells. Alternatively antibodies able to bind surface immunoglobulin and other molecules on B cells as well as antibodies able to bind the TCR and other molecules on T cells can be combined with the pharmaceutical composition of the invention.

The pharmaceutical composition of the invention may be in the form of a liposome in which protein of the present invention is combined, in addition to other pharmaceutically acceptable carriers, with amphipathic agents such as lipids which exist in aggregated form as micelles, insoluble monolayers, liquid crystals, or lamellar layers in aqueous solution. Suitable lipids for liposomal formulation include, without limitation, monoglycerides, diglycerides, sulfatides, lysolecithins, phospholipids, saponin, bile acids, and the like. Preparation of such liposomal formulations is within the level of skill in the art, as disclosed, for example, in U.S. Patent Nos. 4,235,871; 4,501,728; 4,837,028; and 4,737,323, all of which are incorporated herein by reference.

The amount of protein or other active ingredient of the present invention in the pharmaceutical composition of the present invention will depend upon the nature and severity of the condition being treated, and on the nature of prior treatments which the patient has undergone. Ultimately, the attending physician will decide the amount of protein or other active ingredient of the present invention with which to treat each individual patient. Initially, the attending physician will administer low doses of protein or other active ingredient of the present

invention and observe the patient's response. Larger doses of protein or other active ingredient of the present invention may be administered until the optimal therapeutic effect is obtained for the patient, and at that point the dosage is not increased further. It is contemplated that the various pharmaceutical compositions used to practice the method of the present invention should contain about 0.01 µg to about 100 mg (preferably about 0.1 µg to about 10 mg, more preferably about 0.1 µg to about 1 mg) of protein or other active ingredient of the present invention per kg body weight. For compositions of the present invention which are useful for bone, cartilage, tendon or ligament regeneration, the therapeutic method includes administering the composition topically, systematically, or locally as an implant or device. When administered, the therapeutic composition for use in this invention is, of course, in a pyrogen-free, physiologically acceptable form. Further, the composition may desirably be encapsulated or injected in a viscous form for delivery to the site of bone, cartilage or tissue damage. Topical administration may be suitable for wound healing and tissue repair. Therapeutically useful agents other than a protein or other active ingredient of the invention which may also optionally be included in the composition as described above, may alternatively or additionally, be administered simultaneously or sequentially with the composition in the methods of the invention. Preferably for bone and/or cartilage formation, the composition would include a matrix capable of delivering the protein-containing or other active ingredient-containing composition to the site of bone and/or cartilage damage, providing a structure for the developing bone and cartilage and optimally capable of being resorbed into the body. Such matrices may be formed of materials presently in use for other implanted medical applications.

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The choice of matrix material is based on biocompatibility, biodegradability, mechanical properties, cosmetic appearance and interface properties. The particular application of the compositions will define the appropriate formulation. Potential matrices for the compositions may be biodegradable and chemically defined calcium sulfate, tricalcium phosphate, hydroxyapatite, polylactic acid, polyglycolic acid and polyanhydrides. Other potential materials are biodegradable and biologically well-defined, such as bone or dermal collagen. Further matrices are comprised of pure proteins or extracellular matrix components. Other potential matrices are nonbiodegradable and chemically defined, such as sintered hydroxyapatite, bioglass, aluminates, or other ceramics. Matrices may be comprised of combinations of any of the above mentioned types of material, such as polylactic acid and hydroxyapatite or collagen and tricalcium phosphate. The bioceramics may be altered in composition, such as in calcium-aluminate-phosphate and processing to alter pore size, particle size, particle shape, and biodegradability. Presently preferred is a 50:50 (mole weight) copolymer of lactic acid and glycolic acid in the form of porous particles having diameters ranging from 150 to 800 microns.

In some applications, it will be useful to utilize a sequestering agent, such as carboxymethyl cellulose or autologous blood clot, to prevent the protein compositions from disassociating from the matrix.

A preferred family of sequestering agents is cellulosic materials such as alkylcelluloses (including hydroxyalkylcelluloses), including methylcellulose, ethylcellulose, hydroxyethylcellulose, hydroxypropylcellulose, hydroxypropyl-methylcellulose, and carboxymethylcellulose, the most preferred being cationic salts of carboxymethylcellulose (CMC). Other preferred sequestering agents include hyaluronic acid, sodium alginate, poly(ethylene glycol), polyoxyethylene oxide, carboxyvinyl polymer and poly(vinyl alcohol). The amount of sequestering agent useful herein is 0.5-20 wt %, preferably 1-10 wt % based on total formulation weight, which represents the amount necessary to prevent desorption of the protein from the polymer matrix and to provide appropriate handling of the composition, yet not so much that the progenitor cells are prevented from infiltrating the matrix, thereby providing the protein the opportunity to assist the osteogenic activity of the progenitor cells. In further compositions, proteins or other active ingredients of the invention may be combined with other agents beneficial to the treatment of the bone and/or cartilage defect, wound, or tissue in question. These agents include various growth factors such as epidermal growth factor (EGF), platelet derived growth factor (PDGF), transforming growth factors (TGF-α and TGF-β), and insulin-like growth factor (IGF).

The therapeutic compositions are also presently valuable for veterinary applications. Particularly domestic animals and thoroughbred horses, in addition to humans, are desired patients for such treatment with proteins or other active ingredients of the present invention. The dosage regimen of a protein-containing pharmaceutical composition to be used in tissue regeneration will be determined by the attending physician considering various factors which modify the action of the proteins, *e.g.*, amount of tissue weight desired to be formed, the site of damage, the condition of the damaged tissue, the size of a wound, type of damaged tissue (*e.g.*, bone), the patient's age, sex, and diet, the severity of any infection, time of administration and other clinical factors. The dosage may vary with the type of matrix used in the reconstitution and with inclusion of other proteins in the pharmaceutical composition. For example, the addition of other known growth factors, such as IGF I (insulin like growth factor I), to the final composition, may also effect the dosage. Progress can be monitored by periodic assessment of tissue/bone growth and/or repair, for example, X-rays, histomorphometric determinations and tetracycline labeling.

Polynucleotides of the present invention can also be used for gene therapy. Such polynucleotides can be introduced either in vivo or ex vivo into cells for expression in a

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mammalian subject. Polynucleotides of the invention may also be administered by other known methods for introduction of nucleic acid into a cell or organism (including, without limitation, in the form of viral vectors or naked DNA). Cells may also be cultured ex vivo in the presence of proteins of the present invention in order to proliferate or to produce a desired effect on or activity in such cells. Treated cells can then be introduced in vivo for therapeutic purposes.

4.12.3 EFFECTIVE DOSAGE

Pharmaceutical compositions suitable for use in the present invention include compositions wherein the active ingredients are contained in an effective amount to achieve its intended purpose. More specifically, a therapeutically effective amount means an amount effective to prevent development of or to alleviate the existing symptoms of the subject being treated. Determination of the effective amount is well within the capability of those skilled in the art, especially in light of the detailed disclosure provided herein. For any compound used in the method of the invention, the therapeutically effective dose can be estimated initially from appropriate in vitro assays. For example, a dose can be formulated in animal models to achieve a circulating concentration range that can be used to more accurately determine useful doses in humans. For example, a dose can be formulated in animal models to achieve a circulating concentration range that includes the IC₅₀ as determined in cell culture (*i.e.*, the concentration of the test compound which achieves a half-maximal inhibition of the protein's biological activity). Such information can be used to more accurately determine useful doses in humans.

A therapeutically effective dose refers to that amount of the compound that results in amelioration of symptoms or a prolongation of survival in a patient. Toxicity and therapeutic efficacy of such compounds can be determined by standard pharmaceutical procedures in cell cultures or experimental animals, e.g., for determining the LD₅₀ (the dose lethal to 50% of the population) and the ED₅₀ (the dose therapeutically effective in 50% of the population). The dose ratio between toxic and therapeutic effects is the therapeutic index and it can be expressed as the ratio between LD₅₀ and ED₅₀. Compounds which exhibit high therapeutic indices are preferred. The data obtained from these cell culture assays and animal studies can be used in formulating a range of dosage for use in human. The dosage of such compounds lies preferably within a range of circulating concentrations that include the ED₅₀ with little or no toxicity. The dosage may vary within this range depending upon the dosage form employed and the route of administration utilized. The exact formulation, route of administration and dosage can be chosen by the individual physician in view of the patient's condition. See, e.g., Fingl et al., 1975, in "The Pharmacological Basis of Therapeutics", Ch. 1 p.1. Dosage amount and interval may be adjusted individually to provide plasma levels of the active moiety which are sufficient to maintain the

desired effects, or minimal effective concentration (MEC). The MEC will vary for each compound but can be estimated from *in vitro* data. Dosages necessary to achieve the MEC will depend on individual characteristics and route of administration. However, HPLC assays or bioassays can be used to determine plasma concentrations.

Dosage intervals can also be determined using MEC value. Compounds should be administered using a regimen which maintains plasma levels above the MEC for 10-90% of the time, preferably between 30-90% and most preferably between 50-90%. In cases of local administration or selective uptake, the effective local concentration of the drug may not be related to plasma concentration.

An exemplary dosage regimen for polypeptides or other compositions of the invention will be in the range of about $0.01~\mu g/kg$ to 100~mg/kg of body weight daily, with the preferred dose being about $0.1~\mu g/kg$ to 25~mg/kg of patient body weight daily, varying in adults and children. Dosing may be once daily, or equivalent doses may be delivered at longer or shorter intervals.

The amount of composition administered will, of course, be dependent on the subject being treated, on the subject's age and weight, the severity of the affliction, the manner of administration and the judgment of the prescribing physician.

4.12.4 PACKAGING

The compositions may, if desired, be presented in a pack or dispenser device which may contain one or more unit dosage forms containing the active ingredient. The pack may, for example, comprise metal or plastic foil, such as a blister pack. The pack or dispenser device may be accompanied by instructions for administration. Compositions comprising a compound of the invention formulated in a compatible pharmaceutical carrier may also be prepared, placed in an appropriate container, and labeled for treatment of an indicated condition.

4.13 ANTIBODIES

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Also included in the invention are antibodies to proteins, or fragments of proteins of the invention. The term "antibody" as used herein refers to immunoglobulin molecules and immunologically active portions of immunoglobulin (Ig) molecules, i.e., molecules that contain an antigen binding site that specifically binds (immunoreacts with) an antigen. Such antibodies include, but are not limited to, polyclonal, monoclonal, chimeric, single chain, F_{ab} , F_{ab} and $F_{(ab')2}$ fragments, and an F_{ab} expression library. In general, an antibody molecule obtained from humans relates to any of the classes IgG, IgM, IgA, IgE and IgD, which differ from one another by the nature of the heavy chain present in the molecule. Certain classes have subclasses as well,

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such as IgG₁, IgG₂, and others. Furthermore, in humans, the light chain may be a kappa chain or a lambda chain. Reference herein to antibodies includes a reference to all such classes, subclasses and types of human antibody species.

An isolated related protein of the invention may be intended to serve as an antigen, or a portion or fragment thereof, and additionally can be used as an immunogen to generate antibodies that immunospecifically bind the antigen, using standard techniques for polyclonal and monoclonal antibody preparation. The full-length protein can be used or, alternatively, the invention provides antigenic peptide fragments of the antigen for use as immunogens. An antigenic peptide fragment comprises at least 6 amino acid residues of the amino acid sequence of the full length protein, such as an amino acid sequence shown in SEQ ID NO:985, and encompasses an epitope thereof such that an antibody raised against the peptide forms a specific immune complex with the full length protein or with any fragment that contains the epitope. Preferably, the antigenic peptide comprises at least 10 amino acid residues, or at least 15 amino acid residues, or at least 20 amino acid residues, or at least 30 amino acid residues. Preferred epitopes encompassed by the antigenic peptide are regions of the protein that are located on its surface; commonly these are hydrophilic regions.

In certain embodiments of the invention, at least one epitope encompassed by the antigenic peptide is a region of -related protein that is located on the surface of the protein, e.g., a hydrophilic region. A hydrophobicity analysis of the human related protein sequence will indicate which regions of a related protein are particularly hydrophilic and, therefore, are likely to encode surface residues useful for targeting antibody production. As a means for targeting antibody production, hydropathy plots showing regions of hydrophilicity and hydrophobicity may be generated by any method well known in the art, including, for example, the Kyte Doolittle or the Hopp Woods methods, either with or without Fourier transformation. See, e.g., Hopp and Woods, 1981, Proc. Nat. Acad. Sci. USA 78: 3824-3828; Kyte and Doolittle 1982, J. Mol. Biol. 157: 105-142, each of which is incorporated herein by reference in its entirety. Antibodies that are specific for one or more domains within an antigenic protein, or derivatives, fragments, analogs or homologs thereof, are also provided herein.

A protein of the invention, or a derivative, fragment, analog, homolog or ortholog thereof, may be utilized as an immunogen in the generation of antibodies that immunospecifically bind these protein components.

Various procedures known within the art may be used for the production of polyclonal or monoclonal antibodies directed against a protein of the invention, or against derivatives, fragments, analogs homologs or orthologs thereof (see, for example, Antibodies: A Laboratory

Manual, Harlow E, and Lane D, 1988, Cold Spring Harbor Laboratory Press, Cold Spring Harbor, NY, incorporated herein by reference). Some of these antibodies are discussed below.

5.13.1 Polyclonal Antibodies

For the production of polyclonal antibodies, various suitable host animals (e.g., rabbit, goat, mouse or other mammal) may be immunized by one or more injections with the native protein, a synthetic variant thereof, or a derivative of the foregoing. An appropriate immunogenic preparation can contain, for example, the naturally occurring immunogenic protein, a chemically synthesized polypeptide representing the immunogenic protein, or a recombinantly expressed immunogenic protein. Furthermore, the protein may be conjugated to a second protein known to be immunogenic in the mammal being immunized. Examples of such immunogenic proteins include but are not limited to keyhole limpet hemocyanin, serum albumin, bovine thyroglobulin, and soybean trypsin inhibitor. The preparation can further include an adjuvant. Various adjuvants used to increase the immunological response include, but are not limited to, Freund's (complete and incomplete), mineral gels (e.g., aluminum hydroxide), surface active substances (e.g., lysolecithin, pluronic polyols, polyanions, peptides, oil emulsions, dinitrophenol, etc.), adjuvants usable in humans such as Bacille Calmette-Guerin and Corynebacterium parvum, or similar immunostimulatory agents. Additional examples of adjuvants which can be employed include MPL-TDM adjuvant (monophosphoryl Lipid A, synthetic trehalose dicorynomycolate).

The polyclonal antibody molecules directed against the immunogenic protein can be isolated from the mammal (e.g., from the blood) and further purified by well known techniques, such as affinity chromatography using protein A or protein G, which provide primarily the IgG fraction of immune serum. Subsequently, or alternatively, the specific antigen which is the target of the immunoglobulin sought, or an epitope thereof, may be immobilized on a column to purify the immune specific antibody by immunoaffinity chromatography. Purification of immunoglobulins is discussed, for example, by D. Wilkinson (The Scientist, published by The Scientist, Inc., Philadelphia PA, Vol. 14, No. 8 (April 17, 2000), pp. 25-28).

5.13.2 Monoclonal Antibodies

The term "monoclonal antibody" (MAb) or "monoclonal antibody composition", as used herein, refers to a population of antibody molecules that contain only one molecular species of antibody molecule consisting of a unique light chain gene product and a unique heavy chain gene product. In particular, the complementarity determining regions (CDRs) of the monoclonal antibody are identical in all the molecules of the population. MAbs thus contain an antigen

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binding site capable of immunoreacting with a particular epitope of the antigen characterized by a unique binding affinity for it.

Monoclonal antibodies can be prepared using hybridoma methods, such as those described by Kohler and Milstein, Nature, 256:495 (1975). In a hybridoma method, a mouse, hamster, or other appropriate host animal, is typically immunized with an immunizing agent to elicit lymphocytes that produce or are capable of producing antibodies that will specifically bind to the immunizing agent. Alternatively, the lymphocytes can be immunized in vitro. The immunizing agent will typically include the protein antigen, a fragment thereof or a fusion protein thereof. Generally, either peripheral blood lymphocytes are used if cells of human origin are desired, or spleen cells or lymph node cells are used if non-human mammalian sources are desired. The lymphocytes are then fused with an immortalized cell line using a suitable fusing agent, such as polyethylene glycol, to form a hybridoma cell (Goding, Monoclonal Antibodies: Principles and Practice, Academic Press, (1986) pp. 59-103). Immortalized cell lines are usually transformed mammalian cells, particularly myeloma cells of rodent, bovine and human origin. Usually, rat or mouse myeloma cell lines are employed. The hybridoma cells can be cultured in a suitable culture medium that preferably contains one or more substances that inhibit the growth or survival of the unfused, immortalized cells. For example, if the parental cells lack the enzyme hypoxanthine guanine phosphoribosyl transferase (HGPRT or HPRT), the culture medium for the hybridomas typically will include hypoxanthine, aminopterin, and thymidine ("HAT medium"), which substances prevent the growth of HGPRT-deficient cells.

Preferred immortalized cell lines are those that fuse efficiently, support stable high level expression of antibody by the selected antibody-producing cells, and are sensitive to a medium such as HAT medium. More preferred immortalized cell lines are murine myeloma lines, which can be obtained, for instance, from the Salk Institute Cell Distribution Center, San Diego, California and the American Type Culture Collection, Manassas, Virginia. Human myeloma and mouse-human heteromyeloma cell lines also have been described for the production of human monoclonal antibodies (Kozbor, J. Immunol., 133:3001 (1984); Brodeur et al., Monoclonal Antibody Production Techniques and Applications, Marcel Dekker, Inc., New York, (1987) pp. 51-63).

The culture medium in which the hybridoma cells are cultured can then be assayed for the presence of monoclonal antibodies directed against the antigen. Preferably, the binding specificity of monoclonal antibodies produced by the hybridoma cells is determined by immunoprecipitation or by an in vitro binding assay, such as radioimmunoassay (RIA) or enzyme-linked immunoabsorbent assay (ELISA). Such techniques and assays are known in the art. The binding affinity of the monoclonal antibody can, for example, be determined by the

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Scatchard analysis of Munson and Pollard, <u>Anal. Biochem.</u>, <u>107</u>:220 (1980). Preferably, antibodies having a high degree of specificity and a high binding affinity for the target antigen

are isolated.

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After the desired hybridoma cells are identified, the clones can be subcloned by limiting dilution procedures and grown by standard methods. Suitable culture media for this purpose include, for example, Dulbecco's Modified Eagle's Medium and RPMI-1640 medium. Alternatively, the hybridoma cells can be grown in vivo as ascites in a mammal. The monoclonal antibodies secreted by the subclones can be isolated or purified from the culture medium or ascites fluid by conventional immunoglobulin purification procedures such as, for example, protein A-Sepharose, hydroxylapatite chromatography, gel electrophoresis, dialysis, or affinity chromatography.

The monoclonal antibodies can also be made by recombinant DNA methods, such as those described in U.S. Patent No. 4,816,567. DNA encoding the monoclonal antibodies of the invention can be readily isolated and sequenced using conventional procedures (e.g., by using oligonucleotide probes that are capable of binding specifically to genes encoding the heavy and light chains of murine antibodies). The hybridoma cells of the invention serve as a preferred source of such DNA. Once isolated, the DNA can be placed into expression vectors, which are then transfected into host cells such as simian COS cells, Chinese hamster ovary (CHO) cells, or myeloma cells that do not otherwise produce immunoglobulin protein, to obtain the synthesis of monoclonal antibodies in the recombinant host cells. The DNA also can be modified, for example, by substituting the coding sequence for human heavy and light chain constant domains in place of the homologous murine sequences (U.S. Patent No. 4,816,567; Morrison, Nature 368, 812-13 (1994)) or by covalently joining to the immunoglobulin coding sequence all or part of the coding sequence for a non-immunoglobulin polypeptide. Such a non-immunoglobulin polypeptide can be substituted for the constant domains of an antibody of the invention, or can be substituted for the variable domains of one antigen-combining site of an antibody of the invention to create a chimeric bivalent antibody.

5.13.2 Humanized Antibodies

The antibodies directed against the protein antigens of the invention can further comprise humanized antibodies or human antibodies. These antibodies are suitable for administration to humans without engendering an immune response by the human against the administered immunoglobulin. Humanized forms of antibodies are chimeric immunoglobulins, immunoglobulin chains or fragments thereof (such as Fv, Fab, Fab', F(ab')₂ or other antigenbinding subsequences of antibodies) that are principally comprised of the sequence of a human

immunoglobulin, and contain minimal sequence derived from a non-human immunoglobulin. Humanization can be performed following the method of Winter and co-workers (Jones et al., Nature, 321:522-525 (1986); Riechmann et al., Nature, 332:323-327 (1988); Verhoeyen et al., Science, 239:1534-1536 (1988)), by substituting rodent CDRs or CDR sequences for the corresponding sequences of a human antibody. (See also U.S. Patent No. 5,225,539.) In some instances, Fv framework residues of the human immunoglobulin are replaced by corresponding non-human residues. Humanized antibodies can also comprise residues which are found neither in the recipient antibody nor in the imported CDR or framework sequences. In general, the humanized antibody will comprise substantially all of at least one, and typically two, variable domains, in which all or substantially all of the CDR regions correspond to those of a non-human immunoglobulin and all or substantially all of the framework regions are those of a human immunoglobulin consensus sequence. The humanized antibody optimally also will comprise at least a portion of an immunoglobulin constant region (Fc), typically that of a human immunoglobulin (Jones et al., 1986; Riechmann et al., 1988; and Presta, Curr. Op. Struct. Biol., 2:593-596 (1992)).

5.13.3 Human Antibodies

Fully human antibodies relate to antibody molecules in which essentially the entire sequences of both the light chain and the heavy chain, including the CDRs, arise from human genes. Such antibodies are termed "human antibodies", or "fully human antibodies" herein. Human monoclonal antibodies can be prepared by the trioma technique; the human B-cell hybridoma technique (see Kozbor, et al., 1983 Immunol Today 4: 72) and the EBV hybridoma technique to produce human monoclonal antibodies (see Cole, et al., 1985 In: MONOCLONAL ANTIBODIES AND CANCER THERAPY, Alan R. Liss, Inc., pp. 77-96). Human monoclonal antibodies may be utilized in the practice of the present invention and may be produced by using human hybridomas (see Cote, et al., 1983. Proc Natl Acad Sci USA 80: 2026-2030) or by transforming human B-cells with Epstein Barr Virus in vitro (see Cole, et al., 1985 In: MONOCLONAL ANTIBODIES AND CANCER THERAPY, Alan R. Liss, Inc., pp. 77-96).

In addition, human antibodies can also be produced using additional techniques, including phage display libraries (Hoogenboom and Winter, J. Mol. Biol., 227:381 (1991); Marks et al., J. Mol. Biol., 222:581 (1991)). Similarly, human antibodies can be made by introducing human immunoglobulin loci into transgenic animals, e.g., mice in which the endogenous immunoglobulin genes have been partially or completely inactivated. Upon challenge, human antibody production is observed, which closely resembles that seen in humans in all respects, including gene rearrangement, assembly, and antibody repertoire. This approach

is described, for example, in U.S. Patent Nos. 5,545,807; 5,545,806; 5,569,825; 5,625,126; 5,633,425; 5,661,016, and in Marks et al. (Bio/Technology 10, 779-783 (1992)); Lonberg et al. (Nature 368 856-859 (1994)); Morrison (Nature 368, 812-13 (1994)); Fishwild et al., (Nature Biotechnology 14, 845-51 (1996)); Neuberger (Nature Biotechnology 14, 826 (1996)); and Lonberg and Huszar (Intern. Rev. Immunol. 13 65-93 (1995)).

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Human antibodies may additionally be produced using transgenic nonhuman animals which are modified so as to produce fully human antibodies rather than the animal's endogenous antibodies in response to challenge by an antigen. (See PCT publication WO94/02602). The endogenous genes encoding the heavy and light immunoglobulin chains in the nonhuman host have been incapacitated, and active loci encoding human heavy and light chain immunoglobulins are inserted into the host's genome. The human genes are incorporated, for example, using yeast artificial chromosomes containing the requisite human DNA segments. An animal which provides all the desired modifications is then obtained as progeny by crossbreeding intermediate transgenic animals containing fewer than the full complement of the modifications. The preferred embodiment of such a nonhuman animal is a mouse, and is termed the XenomouseTM as disclosed in PCT publications WO 96/33735 and WO 96/34096. This animal produces B cells which secrete fully human immunoglobulins. The antibodies can be obtained directly from the animal after immunization with an immunogen of interest, as, for example, a preparation of a polyclonal antibody, or alternatively from immortalized B cells derived from the animal, such as hybridomas producing monoclonal antibodies. Additionally, the genes encoding the immunoglobulins with human variable regions can be recovered and expressed to obtain the antibodies directly, or can be further modified to obtain analogs of antibodies such as, for example, single chain Fv molecules.

An example of a method of producing a nonhuman host, exemplified as a mouse, lacking expression of an endogenous immunoglobulin heavy chain is disclosed in U.S. Patent No. 5,939,598. It can be obtained by a method including deleting the J segment genes from at least one endogenous heavy chain locus in an embryonic stem cell to prevent rearrangement of the locus and to prevent formation of a transcript of a rearranged immunoglobulin heavy chain locus, the deletion being effected by a targeting vector containing a gene encoding a selectable marker; and producing from the embryonic stem cell a transgenic mouse whose somatic and germ cells contain the gene encoding the selectable marker.

A method for producing an antibody of interest, such as a human antibody, is disclosed in U.S. Patent No. 5,916,771. It includes introducing an expression vector that contains a nucleotide sequence encoding a heavy chain into one mammalian host cell in culture, introducing an expression vector containing a nucleotide sequence encoding a light chain into another

mammalian host cell, and fusing the two cells to form a hybrid cell. The hybrid cell expresses an antibody containing the heavy chain and the light chain.

In a further improvement on this procedure, a method for identifying a clinically relevant epitope on an immunogen, and a correlative method for selecting an antibody that binds immunospecifically to the relevant epitope with high affinity, are disclosed in PCT publication WO 99/53049.

5.13.4 Fab Fragments and Single Chain Antibodies

According to the invention, techniques can be adapted for the production of single-chain antibodies specific to an antigenic protein of the invention (see e.g., U.S. Patent No. 4,946,778). In addition, methods can be adapted for the construction of F_{ab} expression libraries (see e.g., Huse, et al., 1989 Science 246: 1275-1281) to allow rapid and effective identification of monoclonal F_{ab} fragments with the desired specificity for a protein or derivatives, fragments, analogs or homologs thereof. Antibody fragments that contain the idiotypes to a protein antigen may be produced by techniques known in the art including, but not limited to: (i) an $F_{(ab)2}$ fragment produced by pepsin digestion of an antibody molecule; (ii) an F_{ab} fragment generated by reducing the disulfide bridges of an $F_{(ab)2}$ fragment; (iii) an F_{ab} fragment generated by the treatment of the antibody molecule with papain and a reducing agent and (iv) F_{v} fragments.

5.13.5 Bispecific Antibodies

Bispecific antibodies are monoclonal, preferably human or humanized, antibodies that have binding specificities for at least two different antigens. In the present case, one of the binding specificities is for an antigenic protein of the invention. The second binding target is any other antigen, and advantageously is a cell-surface protein or receptor or receptor subunit.

Methods for making bispecific antibodies are known in the art. Traditionally, the recombinant production of bispecific antibodies is based on the co-expression of two immunoglobulin heavy-chain/light-chain pairs, where the two heavy chains have different specificities (Milstein and Cuello, Nature, 305:537-539 (1983)). Because of the random assortment of immunoglobulin heavy and light chains, these hybridomas (quadromas) produce a potential mixture of ten different antibody molecules, of which only one has the correct bispecific structure. The purification of the correct molecule is usually accomplished by affinity chromatography steps. Similar procedures are disclosed in WO 93/08829, published 13 May 1993, and in Traunecker *et al.*, 1991 *EMBO J.*, 10:3655-3659.

Antibody variable domains with the desired binding specificities (antibody-antigen combining sites) can be fused to immunoglobulin constant domain sequences. The fusion

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preferably is with an immunoglobulin heavy-chain constant domain, comprising at least part of the hinge, CH2, and CH3 regions. It is preferred to have the first heavy-chain constant region (CH1) containing the site necessary for light-chain binding present in at least one of the fusions. DNAs encoding the immunoglobulin heavy-chain fusions and, if desired, the immunoglobulin light chain, are inserted into separate expression vectors, and are co-transfected into a suitable host organism. For further details of generating bispecific antibodies see, for example, Suresh et al., Methods in Enzymology, 121:210 (1986).

According to another approach described in WO 96/27011, the interface between a pair of antibody molecules can be engineered to maximize the percentage of heterodimers which are recovered from recombinant cell culture. The preferred interface comprises at least a part of the CH3 region of an antibody constant domain. In this method, one or more small amino acid side chains from the interface of the first antibody molecule are replaced with larger side chains (e.g. tyrosine or tryptophan). Compensatory "cavities" of identical or similar size to the large side chain(s) are created on the interface of the second antibody molecule by replacing large amino acid side chains with smaller ones (e.g. alanine or threonine). This provides a mechanism for increasing the yield of the heterodimer over other unwanted end-products such as homodimers.

Bispecific antibodies can be prepared as full length antibodies or antibody fragments (e.g. F(ab')₂ bispecific antibodies). Techniques for generating bispecific antibodies from antibody fragments have been described in the literature. For example, bispecific antibodies can be prepared using chemical linkage. Brennan et al., Science 229:81 (1985) describe a procedure wherein intact antibodies are proteolytically cleaved to generate F(ab')₂ fragments. These fragments are reduced in the presence of the dithiol complexing agent sodium arsenite to stabilize vicinal dithiols and prevent intermolecular disulfide formation. The Fab' fragments generated are then converted to thionitrobenzoate (TNB) derivatives. One of the Fab'-TNB derivatives is then reconverted to the Fab'-thiol by reduction with mercaptoethylamine and is mixed with an equimolar amount of the other Fab'-TNB derivative to form the bispecific antibody. The bispecific antibodies produced can be used as agents for the selective immobilization of enzymes.

Additionally, Fab' fragments can be directly recovered from E. coli and chemically coupled to form bispecific antibodies. Shalaby et al., <u>J. Exp. Med.</u> 175:217-225 (1992) describe the production of a fully humanized bispecific antibody F(ab')₂ molecule. Each Fab' fragment was separately secreted from E. coli and subjected to directed chemical coupling in vitro to form the bispecific antibody. The bispecific antibody thus formed was able to bind to cells overexpressing the ErbB2 receptor and normal human T cells, as well as trigger the lytic activity of human cytotoxic lymphocytes against human breast tumor targets.

Various techniques for making and isolating bispecific antibody fragments directly from recombinant cell culture have also been described. For example, bispecific antibodies have been produced using leucine zippers. Kostelny et al., J. Immunol. 148(5):1547-1553 (1992). The leucine zipper peptides from the Fos and Jun proteins were linked to the Fab' portions of two different antibodies by gene fusion. The antibody homodimers were reduced at the hinge region to form monomers and then re-oxidized to form the antibody heterodimers. This method can also be utilized for the production of antibody homodimers. The "diabody" technology described by Hollinger et al., Proc. Natl. Acad. Sci. USA 90:6444-6448 (1993) has provided an alternative mechanism for making bispecific antibody fragments. The fragments comprise a heavy-chain variable domain (V_L) to connected to a light-chain variable domain (V_L) by a linker which is too short to allow pairing between the two domains on the same chain. Accordingly, the V_H and V_L domains of one fragment are forced to pair with the complementary V_L and V_H domains of another fragment, thereby forming two antigen-binding sites. Another strategy for making bispecific antibody fragments by the use of single-chain Fv (sFv) dimers has also been reported. See, Gruber et al., J. Immunol. 152:5368 (1994).

Antibodies with more than two valencies are contemplated. For example, trispecific antibodies can be prepared. Tutt et al., J. Immunol. 147:60 (1991). Exemplary bispecific antibodies can bind to two different epitopes, at least one of which originates in the protein antigen of the invention. Alternatively, an anti-antigenic arm of an immunoglobulin molecule can be combined with an arm which binds to a triggering molecule on a leukocyte such as a T-cell receptor molecule (e.g. CD2, CD3, CD28, or B7), or Fc receptors for IgG (FcγR), such as FcγRI (CD64), FcγRII (CD32) and FcγRII (CD16) so as to focus cellular defense mechanisms to the cell expressing the particular antigen. Bispecific antibodies can also be used to direct cytotoxic agents to cells which express a particular antigen. These antibodies possess an antigen-binding arm and an arm which binds a cytotoxic agent or a radionuclide chelator, such as EOTUBE, DPTA, DOTA, or TETA. Another bispecific antibody of interest binds the protein antigen described herein and further binds tissue factor (TF).

5.13.6 Heteroconjugate Antibodies

Heteroconjugate antibodies are also within the scope of the present invention. Heteroconjugate antibodies are composed of two covalently joined antibodies. Such antibodies have, for example, been proposed to target immune system cells to unwanted cells (U.S. Patent No. 4,676,980), and for treatment of HIV infection (WO 91/00360; WO 92/200373; EP 03089). It is contemplated that the antibodies can be prepared in vitro using known methods in synthetic protein chemistry, including those involving crosslinking agents. For example, immunotoxins

can be constructed using a disulfide exchange reaction or by forming a thioether bond. Examples of suitable reagents for this purpose include iminothiolate and methyl-4-mercaptobutyrimidate and those disclosed, for example, in U.S. Patent No. 4,676,980.

5.13.7 Effector Function Engineering

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It can be desirable to modify the antibody of the invention with respect to effector function, so as to enhance, e.g., the effectiveness of the antibody in treating cancer. For example, cysteine residue(s) can be introduced into the Fc region, thereby allowing interchain disulfide bond formation in this region. The homodimeric antibody thus generated can have improved internalization capability and/or increased complement-mediated cell killing and antibody-dependent cellular cytotoxicity (ADCC). See Caron et al., J. Exp Med., 176: 1191-1195 (1992) and Shopes, J. Immunol., 148: 2918-2922 (1992). Homodimeric antibodies with enhanced anti-tumor activity can also be prepared using heterobifunctional cross-linkers as described in Wolff et al. Cancer Research, 53: 2560-2565 (1993). Alternatively, an antibody can be engineered that has dual Fc regions and can thereby have enhanced complement lysis and ADCC capabilities. See Stevenson et al., Anti-Cancer Drug Design, 3: 219-230 (1989).

5.13.8 Immunoconjugates

The invention also pertains to immunoconjugates comprising an antibody conjugated to a cytotoxic agent such as a chemotherapeutic agent, toxin (e.g., an enzymatically active toxin of bacterial, fungal, plant, or animal origin, or fragments thereof), or a radioactive isotope (i.e., a radioconjugate).

Chemotherapeutic agents useful in the generation of such immunoconjugates have been described above. Enzymatically active toxins and fragments thereof that can be used include diphtheria A chain, nonbinding active fragments of diphtheria toxin, exotoxin A chain (from Pseudomonas aeruginosa), ricin A chain, abrin A chain, modeccin A chain, alpha-sarcin, Aleurites fordii proteins, dianthin proteins, Phytolaca americana proteins (PAPI, PAPII, and PAP-S), momordica charantia inhibitor, curcin, crotin, sapaonaria officinalis inhibitor, gelonin, mitogellin, restrictocin, phenomycin, enomycin, and the tricothecenes. A variety of radionuclides are available for the production of radioconjugated antibodies. Examples include ²¹²Bi, ¹³¹I, ¹³¹In, ⁹⁰Y, and ¹⁸⁶Re.

Conjugates of the antibody and cytotoxic agent are made using a variety of bifunctional protein-coupling agents such as N-succinimidyl-3-(2-pyridyldithiol) propionate (SPDP), iminothiolane (IT), bifunctional derivatives of imidoesters (such as dimethyl adipimidate HCL), active esters (such as disuccinimidyl suberate), aldehydes (such as glutareldehyde), bis-azido

compounds (such as bis (p-azidobenzoyl) hexanediamine), bis-diazonium derivatives (such as bis-(p-diazoniumbenzoyl)-ethylenediamine), diisocyanates (such as tolyene 2,6-diisocyanate), and bis-active fluorine compounds (such as 1,5-difluoro-2,4-dinitrobenzene). For example, a ricin immunotoxin can be prepared as described in Vitetta et al., Science, 238: 1098 (1987).

Carbon-14-labeled 1-isothiocyanatobenzyl-3-methyldiethylene triaminepentaacetic acid (MX-DTPA) is an exemplary chelating agent for conjugation of radionucleotide to the antibody. See WO94/11026.

In another embodiment, the antibody can be conjugated to a "receptor" (such streptavidin) for utilization in tumor pretargeting wherein the antibody-receptor conjugate is administered to the patient, followed by removal of unbound conjugate from the circulation using a clearing agent and then administration of a "ligand" (e.g., avidin) that is in turn conjugated to a cytotoxic agent.

4.14 COMPUTER READABLE SEQUENCES

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In one application of this embodiment, a nucleotide sequence of the present invention can be recorded on computer readable media. As used herein, "computer readable media" refers to any medium which can be read and accessed directly by a computer. Such media include, but are not limited to: magnetic storage media, such as floppy discs, hard disc storage medium, and magnetic tape; optical storage media such as CD-ROM; electrical storage media such as RAM and ROM; and hybrids of these categories such as magnetic/optical storage media. A skilled artisan can readily appreciate how any of the presently known computer readable mediums can be used to create a manufacture comprising computer readable medium having recorded thereon a nucleotide sequence of the present invention. As used herein, "recorded" refers to a process for storing information on computer readable medium. A skilled artisan can readily adopt any of the presently known methods for recording information on computer readable medium to generate manufactures comprising the nucleotide sequence information of the present invention.

A variety of data storage structures are available to a skilled artisan for creating a computer readable medium having recorded thereon a nucleotide sequence of the present invention. The choice of the data storage structure will generally be based on the means chosen to access the stored information. In addition, a variety of data processor programs and formats can be used to store the nucleotide sequence information of the present invention on computer readable medium. The sequence information can be represented in a word processing text file, formatted in commercially-available software such as WordPerfect and Microsoft Word, or represented in the form of an ASCII file, stored in a database application, such as DB2, Sybase, Oracle, or the like. A skilled artisan can readily adapt any number of data processor structuring

formats (e.g. text file or database) in order to obtain computer readable medium having recorded thereon the nucleotide sequence information of the present invention.

By providing any of the nucleotide sequences SEQ ID NO: 1-984, 1969-2952, 3937-3942 or 3949-3954 or a representative fragment thereof; or a nucleotide sequence at least 95% identical to any of the nucleotide sequences of SEQ ID NO: 1-984, 1969-2952, 3937-3942 or 3949-3954 in computer readable form, a skilled artisan can routinely access the sequence information for a variety of purposes. Computer software is publicly available which allows a

examples which follow demonstrate how software which implements the BLAST (Altschul et al., J. Mol. Biol. 215:403-410 (1990)) and BLAZE (Brutlag et al., Comp. Chem. 17:203-207

skilled artisan to access sequence information provided in a computer readable medium. The

(1993)) search algorithms on a Sybase system is used to identify open reading frames (ORFs) within a nucleic acid sequence. Such ORFs may be protein encoding fragments and may be

useful in producing commercially important proteins such as enzymes used in fermentation

reactions and in the production of commercially useful metabolites.

As used herein, "a computer-based system" refers to the hardware means, software means, and data storage means used to analyze the nucleotide sequence information of the present invention. The minimum hardware means of the computer-based systems of the present invention comprises a central processing unit (CPU), input means, output means, and data storage means. A skilled artisan can readily appreciate that any one of the currently available computer-based systems are suitable for use in the present invention. As stated above, the computer-based systems of the present invention comprise a data storage means having stored therein a nucleotide sequence of the present invention and the necessary hardware means and software means for supporting and implementing a search means. As used herein, "data storage means" refers to memory which can store nucleotide sequence information of the present invention, or a memory access means which can access manufactures having recorded thereon the nucleotide sequence information of the present invention.

As used herein, "search means" refers to one or more programs which are implemented on the computer-based system to compare a target sequence or target structural motif with the sequence information stored within the data storage means. Search means are used to identify fragments or regions of a known sequence which match a particular target sequence or target motif. A variety of known algorithms are disclosed publicly and a variety of commercially available software for conducting search means are and can be used in the computer-based systems of the present invention. Examples of such software includes, but is not limited to, Smith-Waterman, MacPattern (EMBL), BLASTN and BLASTA (NPOLYPEPTIDEIA). A skilled artisan can readily recognize that any one of the available algorithms or implementing

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software packages for conducting homology searches can be adapted for use in the present computer-based systems. As used herein, a "target sequence" can be any nucleic acid or amino acid sequence of six or more nucleotides or two or more amino acids. A skilled artisan can readily recognize that the longer a target sequence is, the less likely a target sequence will be present as a random occurrence in the database. The most preferred sequence length of a target sequence is from about 10 to 300 amino acids, more preferably from about 30 to 100 nucleotide residues. However, it is well recognized that searches for commercially important fragments, such as sequence fragments involved in gene expression and protein processing, may be of shorter length.

As used herein, "a target structural motif," or "target motif," refers to any rationally selected sequence or combination of sequences in which the sequence(s) are chosen based on a three-dimensional configuration which is formed upon the folding of the target motif. There are a variety of target motifs known in the art. Protein target motifs include, but are not limited to, enzyme active sites and signal sequences. Nucleic acid target motifs include, but are not limited to, promoter sequences, hairpin structures and inducible expression elements (protein binding sequences).

4.15 TRIPLE HELIX FORMATION

In addition, the fragments of the present invention, as broadly described, can be used to control gene expression through triple helix formation or antisense DNA or RNA, both of which methods are based on the binding of a polynucleotide sequence to DNA or RNA.

Polynucleotides suitable for use in these methods are preferably 20 to 40 bases in length and are designed to be complementary to a region of the gene involved in transcription (triple helix - see Lee et al., Nucl. Acids Res. 6:3073 (1979); Cooney et al., Science 15241:456 (1988); and Dervan et al., Science 251:1360 (1991)) or to the mRNA itself (antisense - Olmno, J. Neurochem. 56:560 (1991); Oligodeoxynucleotides as Antisense Inhibitors of Gene Expression, CRC Press, Boca Raton, FL (1988)). Triple helix-formation optimally results in a shut-off of RNA transcription from DNA, while antisense RNA hybridization blocks translation of an mRNA molecule into polypeptide. Both techniques have been demonstrated to be effective in model systems. Information contained in the sequences of the present invention is necessary for the design of an antisense or triple helix oligonucleotide.

4.16 DIAGNOSTIC ASSAYS AND KITS

The present invention further provides methods to identify the presence or expression of one of the ORFs of the present invention, or homolog thereof, in a test sample, using a nucleic

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acid probe or antibodies of the present invention, optionally conjugated or otherwise associated with a suitable label.

In general, methods for detecting a polynucleotide of the invention can comprise contacting a sample with a compound that binds to and forms a complex with the polynucleotide for a period sufficient to form the complex, and detecting the complex, so that if a complex is detected, a polynucleotide of the invention is detected in the sample. Such methods can also comprise contacting a sample under stringent hybridization conditions with nucleic acid primers that anneal to a polynucleotide of the invention under such conditions, and amplifying annealed polynucleotides, so that if a polynucleotide is amplified, a polynucleotide of the invention is detected in the sample.

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In general, methods for detecting a polypeptide of the invention can comprise contacting a sample with a compound that binds to and forms a complex with the polypeptide for a period sufficient to form the complex, and detecting the complex, so that if a complex is detected, a polypeptide of the invention is detected in the sample.

In detail, such methods comprise incubating a test sample with one or more of the antibodies or one or more of the nucleic acid probes of the present invention and assaying for binding of the nucleic acid probes or antibodies to components within the test sample.

Conditions for incubating a nucleic acid probe or antibody with a test sample vary. Incubation conditions depend on the format employed in the assay, the detection methods employed, and the type and nature of the nucleic acid probe or antibody used in the assay. One skilled in the art will recognize that any one of the commonly available hybridization, amplification or immunological assay formats can readily be adapted to employ the nucleic acid probes or antibodies of the present invention. Examples of such assays can be found in Chard, T., An Introduction to Radioimmunoassay and Related Techniques, Elsevier Science Publishers, Amsterdam, The Netherlands (1986); Bullock, G.R. et al., Techniques in Immunocytochemistry, Academic Press, Orlando, FL Vol. 1 (1982), Vol. 2 (1983), Vol. 3 (1985); Tijssen, P., Practice and Theory of immunoassays: Laboratory Techniques in Biochemistry and Molecular Biology, Elsevier Science Publishers, Amsterdam, The Netherlands (1985). The test samples of the present invention include cells, protein or membrane extracts of cells, or biological fluids such as sputum, blood, serum, plasma, or urine. The test sample used in the above-described method will vary based on the assay format, nature of the detection method and the tissues, cells or extracts used as the sample to be assayed. Methods for preparing protein extracts or membrane extracts of cells are well known in the art and can be readily be adapted in order to obtain a sample which is compatible with the system utilized.

In another embodiment of the present invention, kits are provided which contain the necessary reagents to carry out the assays of the present invention. Specifically, the invention provides a compartment kit to receive, in close confinement, one or more containers which comprises: (a) a first container comprising one of the probes or antibodies of the present invention; and (b) one or more other containers comprising one or more of the following: wash reagents, reagents capable of detecting presence of a bound probe or antibody.

In detail, a compartment kit includes any kit in which reagents are contained in separate containers. Such containers include small glass containers, plastic containers or strips of plastic or paper. Such containers allows one to efficiently transfer reagents from one compartment to another compartment such that the samples and reagents are not cross-contaminated, and the agents or solutions of each container can be added in a quantitative fashion from one compartment to another. Such containers will include a container which will accept the test sample, a container which contains the antibodies used in the assay, containers which contain wash reagents (such as phosphate buffered saline, Tris-buffers, etc.), and containers which contain the reagents used to detect the bound antibody or probe. Types of detection reagents include labeled nucleic acid probes, labeled secondary antibodies, or in the alternative, if the primary antibody is labeled, the enzymatic, or antibody binding reagents which are capable of reacting with the labeled antibody. One skilled in the art will readily recognize that the disclosed probes and antibodies of the present invention can be readily incorporated into one of the established kit formats which are well known in the art.

4.17 MEDICAL IMAGING

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The novel polypeptides and binding partners of the invention are useful in medical imaging of sites expressing the molecules of the invention (e.g., where the polypeptide of the invention is involved in the immune response, for imaging sites of inflammation or infection). See, e.g., Kunkel et al., U.S. Pat. NO. 5,413,778. Such methods involve chemical attachment of a labeling or imaging agent, administration of the labeled polypeptide to a subject in a pharmaceutically acceptable carrier, and imaging the labeled polypeptide *in vivo* at the target site.

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4.18 SCREENING ASSAYS

Using the isolated proteins and polynucleotides of the invention, the present invention further provides methods of obtaining and identifying agents which bind to a polypeptide encoded by an ORF corresponding to any of the nucleotide sequences set forth in SEQ ID NO:

1-984, 1969-2952, 3937-3942 or 3949-3954, or bind to a specific domain of the polypeptide encoded by the nucleic acid. In detail, said method comprises the steps of:

- (a) contacting an agent with an isolated protein encoded by an ORF of the present invention, or nucleic acid of the invention; and
 - (b) determining whether the agent binds to said protein or said nucleic acid.

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In general, therefore, such methods for identifying compounds that bind to a polynucleotide of the invention can comprise contacting a compound with a polynucleotide of the invention for a time sufficient to form a polynucleotide/compound complex, and detecting the complex, so that if a polynucleotide/compound complex is detected, a compound that binds to a polynucleotide of the invention is identified.

Likewise, in general, therefore, such methods for identifying compounds that bind to a polypeptide of the invention can comprise contacting a compound with a polypeptide of the invention for a time sufficient to form a polypeptide/compound complex, and detecting the complex, so that if a polypeptide/compound complex is detected, a compound that binds to a polynucleotide of the invention is identified.

Methods for identifying compounds that bind to a polypeptide of the invention can also comprise contacting a compound with a polypeptide of the invention in a cell for a time sufficient to form a polypeptide/compound complex, wherein the complex drives expression of a receptor gene sequence in the cell, and detecting the complex by detecting reporter gene sequence expression, so that if a polypeptide/compound complex is detected, a compound that binds a polypeptide of the invention is identified.

Compounds identified via such methods can include compounds which modulate the activity of a polypeptide of the invention (that is, increase or decrease its activity, relative to activity observed in the absence of the compound). Alternatively, compounds identified via such methods can include compounds which modulate the expression of a polynucleotide of the invention (that is, increase or decrease expression relative to expression levels observed in the absence of the compound). Compounds, such as compounds identified via the methods of the invention, can be tested using standard assays well known to those of skill in the art for their ability to modulate activity/expression.

The agents screened in the above assay can be, but are not limited to, peptides, carbohydrates, vitamin derivatives, or other pharmaceutical agents. The agents can be selected and screened at random or rationally selected or designed using protein modeling techniques.

For random screening, agents such as peptides, carbohydrates, pharmaceutical agents and the like are selected at random and are assayed for their ability to bind to the protein encoded by the ORF of the present invention. Alternatively, agents may be rationally selected or designed.

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As used herein, an agent is said to be "rationally selected or designed" when the agent is chosen based on the configuration of the particular protein. For example, one skilled in the art can readily adapt currently available procedures to generate peptides, pharmaceutical agents and the like, capable of binding to a specific peptide sequence, in order to generate rationally designed antipeptide peptides, for example see Hurby et al., Application of Synthetic Peptides: Antisense Peptides," In Synthetic Peptides, A User's Guide, W.H. Freeman, NY (1992), pp. 289-307, and Kaspczak et al., Biochemistry 28:9230-8 (1989), or pharmaceutical agents, or the like.

In addition to the foregoing, one class of agents of the present invention, as broadly described, can be used to control gene expression through binding to one of the ORFs or EMFs of the present invention. As described above, such agents can be randomly screened or rationally designed/selected. Targeting the ORF or EMF allows a skilled artisan to design sequence specific or element specific agents, modulating the expression of either a single ORF or multiple ORFs which rely on the same EMF for expression control. One class of DNA binding agents are agents which contain base residues which hybridize or form a triple helix formation by binding to DNA or RNA. Such agents can be based on the classic phosphodiester, ribonucleic acid backbone, or can be a variety of sulfhydryl or polymeric derivatives which have base attachment capacity.

Agents suitable for use in these methods preferably contain 20 to 40 bases and are designed to be complementary to a region of the gene involved in transcription (triple helix - see Lee et al., Nucl. Acids Res. 6:3073 (1979); Cooney et al., Science 241:456 (1988); and Dervan et al., Science 251:1360 (1991)) or to the mRNA itself (antisense - Okano, J. Neurochem. 56:560 (1991); Oligodeoxynucleotides as Antisense Inhibitors of Gene Expression, CRC Press, Boca Raton, FL (1988)). Triple helix-formation optimally results in a shut-off of RNA transcription from DNA, while antisense RNA hybridization blocks translation of an mRNA molecule into polypeptide. Both techniques have been demonstrated to be effective in model systems. Information contained in the sequences of the present invention is necessary for the design of an antisense or triple helix oligonucleotide and other DNA binding agents.

Agents which bind to a protein encoded by one of the ORFs of the present invention can be used as a diagnostic agent. Agents which bind to a protein encoded by one of the ORFs of the present invention can be formulated using known techniques to generate a pharmaceutical composition.

4.19 USE OF NUCLEIC ACIDS AS PROBES

Another aspect of the subject invention is to provide for polypeptide-specific nucleic acid hybridization probes capable of hybridizing with naturally occurring nucleotide sequences. The

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hybridization probes of the subject invention may be derived from any of the nucleotide sequences SEQ ID NO: 1-984, 1969-2952, 3937-3942 or 3949-3954. Because the corresponding gene is only expressed in a limited number of tissues, a hybridization probe derived from of any of the nucleotide sequences SEQ ID NO: 1-984, 1969-2952, 3937-3942 or 3949-3954 can be used as an indicator of the presence of RNA of cell type of such a tissue in a sample.

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Any suitable hybridization technique can be employed, such as, for example, in situ hybridization. PCR as described in US Patents Nos. 4,683,195 and 4,965,188 provides additional uses for oligonucleotides based upon the nucleotide sequences. Such probes used in PCR may be of recombinant origin, may be chemically synthesized, or a mixture of both. The probe will comprise a discrete nucleotide sequence for the detection of identical sequences or a degenerate pool of possible sequences for identification of closely related genomic sequences.

Other means for producing specific hybridization probes for nucleic acids include the cloning of nucleic acid sequences into vectors for the production of mRNA probes. Such vectors are known in the art and are commercially available and may be used to synthesize RNA probes in vitro by means of the addition of the appropriate RNA polymerase as T7 or SP6 RNA polymerase and the appropriate radioactively labeled nucleotides. The nucleotide sequences may be used to construct hybridization probes for mapping their respective genomic sequences. The nucleotide sequence provided herein may be mapped to a chromosome or specific regions of a chromosome using well known genetic and/or chromosomal mapping techniques. These techniques include in situ hybridization, linkage analysis against known chromosomal markers, hybridization screening with libraries or flow-sorted chromosomal preparations specific to known chromosomes, and the like. The technique of fluorescent in situ hybridization of chromosome spreads has been described, among other places, in Verma et al (1988) Human Chromosomes: A Manual of Basic Techniques, Pergamon Press, New York NY.

Fluorescent *in situ* hybridization of chromosomal preparations and other physical chromosome mapping techniques may be correlated with additional genetic map data. Examples of genetic map data can be found in the 1994 Genome Issue of Science (265:1981f). Correlation between the location of a nucleic acid on a physical chromosomal map and a specific disease (or predisposition to a specific disease) may help delimit the region of DNA associated with that genetic disease. The nucleotide sequences of the subject invention may be used to detect differences in gene sequences between normal, carrier or affected individuals.

4.20 PREPARATION OF SUPPORT BOUND OLIGONUCLEOTIDES

Oligonucleotides, i.e., small nucleic acid segments, may be readily prepared by, for example, directly synthesizing the oligonucleotide by chemical means, as is commonly practiced using an automated oligonucleotide synthesizer.

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Support bound oligonucleotides may be prepared by any of the methods known to those of skill in the art using any suitable support such as glass, polystyrene or Teflon. One strategy is to precisely spot oligonucleotides synthesized by standard synthesizers. Immobilization can be achieved using passive adsorption (Inouye & Hondo, (1990) J. Clin. Microbiol. 28(6) 1469-72); using UV light (Nagata et al., 1985; Dahlen et al., 1987; Morrissey & Collins, (1989) Mol. Cell Probes 3(2) 189-207) or by covalent binding of base modified DNA (Keller et al., 1988; 1989); all references being specifically incorporated herein.

Another strategy that may be employed is the use of the strong biotin-streptavidin interaction as a linker. For example, Broude *et al.* (1994) Proc. Natl. Acad. Sci. USA 91(8) 3072-6, describe the use of biotinylated probes, although these are duplex probes, that are immobilized on streptavidin-coated magnetic beads. Streptavidin-coated beads may be purchased from Dynal, Oslo. Of course, this same linking chemistry is applicable to coating any surface with streptavidin. Biotinylated probes may be purchased from various sources, such as, e.g., Operon Technologies (Alameda, CA).

Nunc Laboratories (Naperville, IL) is also selling suitable material that could be used. Nunc Laboratories have developed a method by which DNA can be covalently bound to the microwell surface termed Covalink NH. CovaLink NH is a polystyrene surface grafted with secondary amino groups (>NH) that serve as bridge-heads for further covalent coupling. CovaLink Modules may be purchased from Nunc Laboratories. DNA molecules may be bound to CovaLink exclusively at the 5'-end by a phosphoramidate bond, allowing immobilization of more than 1 pmol of DNA (Rasmussen et al., (1991) Anal. Biochem. 198(1) 138-42).

The use of CovaLink NH strips for covalent binding of DNA molecules at the 5'-end has been described (Rasmussen et al., (1991). In this technology, a phosphoramidate bond is employed (Chu et al., (1983) Nucleic Acids Res. 11(8) 6513-29). This is beneficial as immobilization using only a single covalent bond is preferred. The phosphoramidate bond joins the DNA to the CovaLink NH secondary amino groups that are positioned at the end of spacer arms covalently grafted onto the polystyrene surface through a 2 nm long spacer arm. To link an oligonucleotide to CovaLink NH via an phosphoramidate bond, the oligonucleotide terminus must have a 5'-end phosphate group. It is, perhaps, even possible for biotin to be covalently bound to CovaLink and then streptavidin used to bind the probes.

4.21 PREPARATION OF NUCLEIC ACID FRAGMENTS

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The nucleic acids may be obtained from any appropriate source, such as cDNAs, genomic DNA, chromosomal DNA, microdissected chromosome bands, cosmid or YAC inserts, and RNA, including mRNA without any amplification steps. For example, Sambrook *et al.* (1989) describes three protocols for the isolation of high molecular weight DNA from mammalian cells (p. 9.14-9.23).

DNA fragments may be prepared as clones in M13, plasmid or lambda vectors and/or prepared directly from genomic DNA or cDNA by PCR or other amplification methods. Samples may be prepared or dispensed in multiwell plates. About 100-1000 ng of DNA samples may be prepared in 2-500 ml of final volume.

The nucleic acids would then be fragmented by any of the methods known to those of skill in the art including, for example, using restriction enzymes as described at 9.24-9.28 of Sambrook *et al.* (1989), shearing by ultrasound and NaOH treatment.

Low pressure shearing is also appropriate, as described by Schriefer *et al.* (1990) Nucleic Acids Res. 18(24) 7455-6, incorporated herein by reference). In this method, DNA samples are passed through a small French pressure cell at a variety of low to intermediate pressures. A lever device allows controlled application of low to intermediate pressures to the cell. The results of these studies indicate that low-pressure shearing is a useful alternative to sonic and enzymatic DNA fragmentation methods.

One particularly suitable way for fragmenting DNA is contemplated to be that using the two base recognition endonuclease, $Cvi\Pi$, described by Fitzgerald *et al.* (1992) Nucleic Acids Res. 20(14) 3753-62. These authors described an approach for the rapid fragmentation and fractionation of DNA into particular sizes that they contemplated to be suitable for shotgun cloning and sequencing.

The restriction endonuclease CviJI normally cleaves the recognition sequence PuGCPy between the G and C to leave blunt ends. Atypical reaction conditions, which alter the specificity of this enzyme (CviJI**), yield a quasi-random distribution of DNA fragments form the small molecule pUC19 (2688 base pairs). Fitzgerald et al. (1992) quantitatively evaluated the randomness of this fragmentation strategy, using a CviJI** digest of pUC19 that was size fractionated by a rapid gel filtration method and directly ligated, without end repair, to a lac Z minus M13 cloning vector. Sequence analysis of 76 clones showed that CviJI** restricts pyGCPy and PuGCPu, in addition to PuGCPy sites, and that new sequence data is accumulated at a rate consistent with random fragmentation.

As reported in the literature, advantages of this approach compared to sonication and agarose gel fractionation include: smaller amounts of DNA are required (0.2-0.5 ug instead of 2-5

More specifically, the linkage method includes dissolving DNA in water (7.5 ng/ul) and denaturing for 10 min. at 95°C and cooling on ice for 10 min. Ice-cold 0.1 M 1-methylimidazole, pH 7.0 (1-MeIm₇), is then added to a final concentration of 10 mM 1-MeIm₇. A ss DNA solution is then dispensed into CovaLink NH strips (75 ul/well) standing on ice.

Carbodiimide 0.2 M 1-ethyl-3-(3-dimethylaminopropyl)-carbodiimide (EDC), dissolved in 10 mM 1-MeIm₇, is made fresh and 25 ul added per well. The strips are incubated for 5 hours at 50°C. After incubation the strips are washed using, e.g., Nunc-Immuno Wash; first the wells are washed 3 times, then they are soaked with washing solution for 5 min., and finally they are washed 3 times (where in the washing solution is 0.4 N NaOH, 0.25% SDS heated to 50°C).

It is contemplated that a further suitable method for use with the present invention is that described in PCT Patent Application WO 90/03382 (Southern & Maskos), incorporated herein by reference. This method of preparing an oligonucleotide bound to a support involves attaching a nucleoside 3'-reagent through the phosphate group by a covalent phosphodiester link to aliphatic hydroxyl groups carried by the support. The oligonucleotide is then synthesized on the supported nucleoside and protecting groups removed from the synthetic oligonucleotide chain under standard conditions that do not cleave the oligonucleotide from the support. Suitable reagents include nucleoside phosphoramidite and nucleoside hydrogen phosphorate.

An on-chip strategy for the preparation of DNA probe for the preparation of DNA probe arrays may be employed. For example, addressable laser-activated photodeprotection may be employed in the chemical synthesis of oligonucleotides directly on a glass surface, as described by Fodor *et al.* (1991) Science 251(4995) 767-73, incorporated herein by reference. Probes may also be immobilized on nylon supports as described by Van Ness *et al.* (1991) Nucleic Acids Res. 19(12) 3345-50; or linked to Teflon using the method of Duncan & Cavalier (1988) Anal. Biochem. 169(1) 104-8; all references being specifically incorporated herein.

To link an oligonucleotide to a nylon support, as described by Van Ness *et al.* (1991), requires activation of the nylon surface via alkylation and selective activation of the 5'-amine of oligonucleotides with cyanuric chloride.

One particular way to prepare support bound oligonucleotides is to utilize the light-generated synthesis described by Pease *et al.*, (1994) PNAS USA 91(11) 5022-6, incorporated herein by reference). These authors used current photolithographic techniques to generate arrays of immobilized oligonucleotide probes (DNA chips). These methods, in which light is used to direct the synthesis of oligonucleotide probes in high-density, miniaturized arrays, utilize photolabile 5'-protected *N*-acyl-deoxynucleoside phosphoramidites, surface linker chemistry and versatile combinatorial synthesis strategies. A matrix of 256 spatially defined oligonucleotide probes may be generated in this manner.

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ug); and fewer steps are involved (no preligation, end repair, chemical extraction, or agarose gel electrophoresis and elution are needed

Irrespective of the manner in which the nucleic acid fragments are obtained or prepared, it is important to denature the DNA to give single stranded pieces available for hybridization. This is achieved by incubating the DNA solution for 2-5 minutes at 80-90°C. The solution is then cooled quickly to 2°C to prevent renaturation of the DNA fragments before they are contacted with the chip. Phosphate groups must also be removed from genomic DNA by methods known in the art.

4.22 PREPARATION OF DNA ARRAYS

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R LANGUAGE=PCLirrays may be prepared by spotting DNA samples on a support such as a nylon membrane.

Using eMatrix software package (Stanford University, Stanford, CA) (Wu et al., J. Comp. Biol., Vol. 6 pp. 219-235 (1999) herein incorporated by reference), all the sequences were examined to determine whether they had identifiable signature regions. Table 3 shows the signature region found in the indicated polypeptide sequences, the description of the signature, the eMatrix p-value(s) and the position(s) of the signature within the polypeptide sequence.

Using the pFam software program (Sonnhammer et al., Nucleic Acids Res., Vol. 26(1) pp. 320-322 (1998) herein incorporated by reference) all the polypeptide sequences were examined for domains with homology to certain peptide domains. Table 4 shows the name of the domain found, the description, the p-value and the pFam score for the identified domain within the sequence.

The nucleotide sequence within the sequences that codes for signal peptide sequences and their cleavage sites can be determine from using Neural Network SignalP V1.1 program (from Center for Biological Sequence Analysis, The Technical University of Denmark). The process for identifying prokaryotic and eukaryotic signal peptides and their cleavage sites are also disclosed by Henrik Nielson, Jacob Engelbrecht, Soren Brunak, and Gunnar von Heijne in the publication "Identification of prokaryotic and eukaryotic signal peptides and prediction of their cleavage sites" Protein Engineering, Vol. 10, no. 1, pp. 1-6 (1997), incorporated herein by reference. A maximum S score and a mean S score, as described in the Nielson et as reference, was obtained for the polypeptide sequences. Table 7 shows the position of the signal peptide in each of the polypeptides and the maximum score and mean score associated with that signal peptide.

5.5 EXAMPLE 5

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Novel Nucleic Acids

Using PHRAP (Univ. of Washington) or CAP4 (Paracel), a full length gene cDNA sequence and its corresponding protein sequence were generated from the assemblage. Any frame shifts and incorrect stop codons were corrected by hand editing. During editing, the sequence was checked using FASTY and/or BLAST against Genbank (i.e., dbEST version 118, gb pri 118, UniGene version 118, Genpept release 118). Other computer programs which may have been used in the editing process were phredPhrap and Consed (University of Washington) and ed-ready, edext and gc-zip-2 (Hyseq, Inc.). The full-length nucleotide, including splice variants resulting from these procedures are shown in the Sequence Listing as SEQ ID NOS: 767-930. The corresponding amino acid sequences are SEQ ID NO:1751-1914.

Table 1 shows the various tissue sources of SEQ ID NO: 767-930.

The homology results for SEQ ID NO: 767-930 were obtained by a BLASTP version 2.0al 19MP-WashU search against Genpept release 120 and Geneseq October 12, 2000 release 21(Derwent), using BLAST algorithm. The nearest neighbor result showed the homologs for SEQ ID NO: 767-930 from Genpept. The translated amino acid sequences for which the nucleic acid sequence encodes are shown in the Sequence Listing. The homologues with identifiable functions for SEQ ID NO: 767-930 are shown in Table 2 below.

Using eMatrix software package (Stanford University, Stanford, CA) (Wu et al., J. Comp. Biol., Vol. 6 pp. 219-235 (1999) herein incorporated by reference), all the sequences were examined to determine whether they had identifiable signature regions. Table 3 shows the signature region found in the indicated polypeptide sequences, the description of the signature, the eMatrix p-value(s) and the position(s) of the signature within the polypeptide sequence.

Using the pFam software program (Sonnhammer et al., Nucleic Acids Res., Vol. 26(1) pp. 320-322 (1998) herein incorporated by reference) all the polypeptide sequences were examined for domains with homology to certain peptide domains. Table 4 shows the name of the domain found, the description, the p-value and the pFam score for the identified domain within the sequence.

The nucleotide sequence within the sequences that codes for signal peptide sequences and their cleavage sites can be determine from using Neural Network SignalP V1.1 program (from Center for Biological Sequence Analysis, The Technical University of Denmark). The process for identifying prokaryotic and eukaryotic signal peptides and their cleavage sites are also disclosed by Henrik Nielson, Jacob Engelbrecht, Soren Brunak, and Gunnar von Heijne in the publication "Identification of prokaryotic and eukaryotic signal peptides and prediction of their cleavage sites" Protein Engineering, Vol. 10, no. 1, pp. 1-6 (1997), incorporated herein by reference. A maximum S score and a mean S score, as described in the Nielson et as reference, was obtained for the polypeptide sequences. Table 7 shows the position of the signal peptide in each of the polypeptides and the maximum score and mean score associated with that signal peptide.

5.6 EXAMPLE 6

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Novel Nucleic Acids

Using PHRAP (Univ. of Washington) or CAP4 (Paracel), a full length gene cDNA sequence and its corresponding protein sequence were generated from the assemblage. Any frame shifts and incorrect stop codons were corrected by hand editing. During editing, the sequence was checked using FASTY and/or BLAST against Genbank (i.e. dbEST version 118, gb pri 118, UniGene version 118, Genpept release 118). Other computer programs which may have been used

in the editing process were phredPhrap and Consed (University of Washington) and ed-ready, edext and gc-zip-2 (Hyseq, Inc.). The full-length nucleotide, including splice variants resulting from these procedures are shown in the Sequence Listing as SEQ ID NOS: 931-965. The corresponding amino acid sequences are shown in SEQ ID NO:1915-1949.

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Table 1 shows the various tissue sources of SEQ ID NO: 931-965.

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The nearest neighbor results for SEQ ID NO: 931-965 were obtained by a BLASTP version 2.0al 19MP-WashU search against Genpept release 120 and Geneseq October 12, 2000 release (Derwent), using BLAST algorithm. The nearest neighbor result showed the closest homologue for SEQ ID NO: 931-965 from Genpept. The translated amino acid sequences for which the nucleic acid sequence encodes are shown in the Sequence Listing. The homologs with identifiable functions for SEQ ID NO: 931-965 are shown in Table 2 below.

Using eMatrix software package (Stanford University, Stanford, CA) (Wu et al., J. Comp. Biol., Vol. 6 pp. 219-235 (1999) herein incorporated by reference), all the sequences were examined to determine whether they had identifiable signature regions. Table 3 shows the signature region found in the indicated polypeptide sequences, the description of the signature, the eMatrix p-value(s) and the position(s) of the signature within the polypeptide sequence.

Using the pFam software program (Sonnhammer et al., Nucleic Acids Res., Vol. 26(1) pp. 320-322 (1998) herein incorporated by reference) all the polypeptide sequences were examined for domains with homology to certain peptide domains. Table 4 shows the name of the domain found, the description, the p-value and the pFam score for the identified domain within the sequence.

The nucleotide sequence within the sequences that codes for signal peptide sequences and their cleavage sites can be determine from using Neural Network SignalP V1.1 program (from Center for Biological Sequence Analysis, The Technical University of Denmark). The process for identifying prokaryotic and eukaryotic signal peptides and their cleavage sites are also disclosed by Henrik Nielson, Jacob Engelbrecht, Soren Brunak, and Gunnar von Heijne in the publication "Identification of prokaryotic and eukaryotic signal peptides and prediction of their cleavage sites" Protein Engineering, Vol. 10, no. 1, pp. 1-6 (1997), incorporated herein by reference. A maximum S score and a mean S score, as described in the Nielson et as reference, was obtained for the polypeptide sequences. Table 7 shows the position of the signal peptide in each of the polypeptides and the maximum score and mean score associated with that signal peptide.

5.7 EXAMPLE 7

Using PHRAP (Univ. of Washington) or CAP4 (Paracel), a full length gene cDNA sequence and its corresponding protein sequence were generated from the assemblage. Any frame shifts and incorrect stop codons were corrected by hand editing. During editing, the sequence was checked using FASTY and/or BLAST against Genbank (i.e. dbEST version 119, gb pri 119, UniGene version 119, Genpept release 119). Other computer programs which may have been used in the editing process were phredPhrap and Consed (University of Washington) and ed-ready, edext and gc-zip-2 (Hyseq, Inc.). The full-length nucleotide, including splice variants resulting from these procedures are shown in the Sequence Listing as SEQ ID NOS:966-974. The corresponding amino acid sequences are SEQ ID NO:1950-1958.

Table 1 shows the various tissue sources of SEQ ID NO: 966-974.

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The nearest neighbor results for SEQ ID NO: 966-974 were obtained by a BLASTP version 2.0al 19MP-WashU search against Genpept release 120 and Geneseq October 12, 2000 release (Derwent), using BLAST algorithm. The nearest neighbor result showed the closest homologue for SEQ ID NO: 966-974 from Genpept. The translated amino acid sequences for which the nucleic acid sequence encodes are shown in the Sequence Listing. The homologs with identifiable functions for SEQ ID NO: 966-974 are shown in Table 2 below.

Using eMatrix software package (Stanford University, Stanford, CA) (Wu et al., J. Comp. Biol., Vol. 6 pp. 219-235 (1999) herein incorporated by reference), all the sequences were examined to determine whether they had identifiable signature regions. Table 3 shows the signature region found in the indicated polypeptide sequences, the description of the signature, the eMatrix p-value(s) and the position(s) of the signature within the polypeptide sequence.

Using the pFam software program (Sonnhammer et al., Nucleic Acids Res., Vol. 26(1) pp. 320-322 (1998) herein incorporated by reference) all the polypeptide sequences were examined for domains with homology to certain peptide domains. Table 4 shows the name of the domain found, the description, the p-value and the pFam score for the identified domain within the sequence.

The nucleotide sequence within the sequences that codes for signal peptide sequences and their cleavage sites can be determine from using Neural Network SignalP V1.1 program (from Center for Biological Sequence Analysis, The Technical University of Denmark). The process for identifying prokaryotic and eukaryotic signal peptides and their cleavage sites are also disclosed by Henrik Nielson, Jacob Engelbrecht, Soren Brunak, and Gunnar von Heijne in the publication "Identification of prokaryotic and eukaryotic signal peptides and prediction of their cleavage sites" Protein Engineering, Vol. 10, no. 1, pp. 1-6 (1997), incorporated herein by reference. A maximum S score and a mean S score, as described in the Nielson et as reference, was obtained for the polypeptide sequences. Table 7 shows the position of the signal peptide in

each of the polypeptides and the maximum score and mean score associated with that signal peptide.

5.8 EXAMPLE 8

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Novel Nucleic Acids

Using PHRAP (Univ. of Washington) or CAP4 (Paracel), a full length gene cDNA sequence and its corresponding protein sequence were generated from the assemblage. Any frame shifts and incorrect stop codons were corrected by hand editing. During editing, the sequence was checked using FASTY and/or BLAST against Genbank (i.e. dbEST version 120, gb pri 120, UniGene version 120, Genpept release 120). Other computer programs which may have been used in the editing process were phredPhrap and Consed (University of Washington) and ed-ready, edext and gc-zip-2 (Hyseq, Inc.). The full-length nucleotide, including splice variants resulting from these procedures are shown in the Sequence Listing as SEQ ID NOS:975-984. The corresponding amino acid sequences are SEQ ID NO:1959-1968.

Table 1 shows the various tissue sources of SEQ ID NO: 975-984.

The nearest neighbor results for SEQ ID NO: 975-984 were obtained by a BLASTP version 2.0al 19MP-WashU search against Genpept release 120 and Geneseq October 21, 2000 release (Derwent), using BLAST algorithm. The nearest neighbor result showed the closest homologue for SEQ ID NO: 975-984 from Genpept. The translated amino acid sequences for which the nucleic acid sequence encodes are shown in the Sequence Listing. The homologs with identifiable functions for SEQ ID NO: 975-984 are shown in Table 2 below.

Using eMatrix software package (Stanford University, Stanford, CA) (Wu et al., J. Comp. Biol., Vol. 6 pp. 219-235 (1999) herein incorporated by reference), all the sequences were examined to determine whether they had identifiable signature regions. Table 3 shows the signature region found in the indicated polypeptide sequences, the description of the signature, the eMatrix p-value(s) and the position(s) of the signature within the polypeptide sequence.

Using the pFam software program (Sonnhammer et al., Nucleic Acids Res., Vol. 26(1) pp. 320-322 (1998) herein incorporated by reference) all the polypeptide sequences were examined for domains with homology to certain peptide domains. Table 4 shows the name of the domain found, the description, the p-value and the pFam score for the identified domain within the sequence.

The nucleotide sequence within the sequences that codes for signal peptide sequences and their cleavage sites can be determine from using Neural Network SignalP V1.1 program (from Center for Biological Sequence Analysis, The Technical University of Denmark). The process for identifying prokaryotic and eukaryotic signal peptides and their cleavage sites are also

disclosed by Henrik Nielson, Jacob Engelbrecht, Soren Brunak, and Gunnar von Heijne in the publication "Identification of prokaryotic and eukaryotic signal peptides and prediction of their cleavage sites" Protein Engineering, Vol. 10, no. 1, pp. 1-6 (1997), incorporated herein by reference. A maximum S score and a mean S score, as described in the Nielson et as reference, was obtained for the polypeptide sequences. Table 7 shows the position of the signal peptide in each of the polypeptides and the maximum score and mean score associated with that signal peptide.

5.9 EXAMPLE 9

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Novel Nucleic Acids

Using PHRAP (Univ. of Washington) or CAP4 (Paracel), a full length gene cDNA sequence and its corresponding protein sequence were generated from the assemblage. Any frame shifts and incorrect stop codons were corrected by hand editing. During editing, the sequence was checked using FASTY and/or BLAST against Genbank (i.e. dbEST version 120, gb pri 120, UniGene version 120, Genpept release 120). Other computer programs which may have been used in the editing process were phredPhrap and Consed (University of Washington) and ed-ready, edext and gc-zip-2 (Hyseq, Inc.). The full-length nucleotide, including splice variants resulting from these procedures are shown in the Sequence Listing as SEQ ID NOS:3937-3942. The corresponding peptide sequence is SEQ ID NO: 3943-3948.

Table 1 shows the various tissue sources of SEQ ID NO: 3937-3942.

The nearest neighbor results for SEQ ID NO: 3937-3942 were obtained by a BLASTP version 2.0al 19MP-WashU search against Genpept release 120 and Geneseq October 12, 2000 release 21 (Derwent), using BLAST algorithm. The nearest neighbor result showed the closest homologue for SEQ ID NO: 3937-3942 from Genpept. The translated amino acid sequences for which the nucleic acid sequence encodes are shown in the Sequence Listing. The homologs with identifiable functions for SEQ ID NO: 3937-3942 are shown in Table 9 below.

Using eMatrix software package (Stanford University, Stanford, CA) (Wu et al., J. Comp. Biol., Vol. 6 pp. 219-235 (1999) herein incorporated by reference), all the sequences were examined to determine whether they had identifiable signature regions. Table 10 shows the signature region found in the indicated polypeptide sequences, the description of the signature, the eMatrix p-value(s) and the position(s) of the signature within the polypeptide sequence.

Using the pFam software program (Sonnhammer et al., Nucleic Acids Res., Vol. 26(1) pp. 320-322 (1998) herein incorporated by reference) all the polypeptide sequences were examined for domains with homology to certain peptide domains. Table 11 shows the name of

the domain found, the description, the p-value and the pFam score for the identified domain within the sequence.

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The nucleotide sequence within the sequences that codes for signal peptide sequences and their cleavage sites can be determine from using Neural Network SignalP V1.1 program (from Center for Biological Sequence Analysis, The Technical University of Denmark). The process for identifying prokaryotic and eukaryotic signal peptides and their cleavage sites are also disclosed by Henrik Nielson, Jacob Engelbrecht, Soren Brunak, and Gunnar von Heijne in the publication "Identification of prokaryotic and eukaryotic signal peptides and prediction of their cleavage sites" Protein Engineering, Vol. 10, no. 1, pp. 1-6 (1997), incorporated herein by reference. A maximum S score and a mean S score, as described in the Nielson et as reference, was obtained for the polypeptide sequences. Table 12 shows the position of the signal peptide in each of the polypeptides and the maximum score and mean score associated with that signal peptide.

Tables 5 and 13 are correlation tables of all of the sequences and the SEQ ID NOS.

TABLE 1

I ID IN A	II ilhmoner	SEQ ID NOS:
i .		
Source	14917116	0 11 05 40 65 55 114 141 156 160 150
		3 11 25 49 65 75 114 141 156 160 172
		190 198 209 217 224 229 234-235 267
		269 274 277 282 284 303 308 312 320
		334 336 352 372 396 398 412 414 437
		453 464 470 481 492-494 508-509 532
		539 581 584 617-619 621 628 633 643
		688 691 745 752 761 768 794 822 837
		848 876 887 953 967 973
GIBCO	AB3001	1 3 12-13 16 22-24 28-29 41 48 58 65 78
		82 89-90 94 97 103 112 114-115 117 120
		122 130-131 168 181 184 186-187 189-
		190 198 208 216 247 249 259 270 277
:		297 301 308 312 314 321 333 348 374
		396 403 406 410 412 416-417 420 423
		426-427 431 456 474 481 484-485 488
		498 500 508-509 530 549 553 558 563-
		564 583 596 602-603 608 612 621-622
		624 643 650 674 699 711 736 738-739
		753 770 779-780 785-786 802-803 816
		822 839 842 848 859 861 871 893-894
		897 900 903 925 954 958 967 969
GIBCO	ABD003	3 19 21-25 28-29 31 33-34 37 39 41 46-48
		53 58 63-64 66 72 78 80 99 103 109-110
		112 114 118 120-124 126 132-133 135
	GIBCO	GIBCO AB3001

			139 143 146 148-149 159 163 168 174
			176 179-180 184-185 188-190 202 208-
			209 216-217 221 223 230 234-235 240
			244 249 251 253 255 258-259 263 269-
			270 277 282 285-286 290 294-295 297
			301-302 304-305 307-308 311-312 314
			320 329 333 335-336 342 344 346 349
			354 358 365 370 373-374 377 380 382-
		ļ	383 388 394-396 399 401-402 406 409-
			410 413 416 420-421 425 428 430-431
			436-437 442 456 462 464 466-467 474
		:	484 486 495-496 500-501 506 508-509
			519 530 537 542 549 561-562 564 572
			574 577-578 580-583 586-587 589 592-
			593 596-597 601 608 610 612-614 617-
			624 630-632 635 637 650 658 663-664
			668 676 679 681 689-690 693 699 724
1			726 732 736 742-743 747 767-770 780
			784 789 793 799 802-805 813 817-818
			822 824 829-831 837 839 845 848 856
			859-860 864 871-872 875-876 881 887
			896-897 901 903 907 910-911 925 930
			933 943-944 947 952-953 958 962-963
		1	965 967 972 977
1 1 1	O1	A DD 001	3 53 66 113 115 126 135 160 172 179 185
adult brain	Clontech	ABR001	204 263 273 305 312 323 358 380 383
			395-396 403 420 428-429 431 461 542
			583 586 606-607 611 620 645-646 688
			690 715 732 736 740 748 754 768 784-
			786 790 796 800 878 897 906-907 947
·			977
adult brain	Clontech	ABR006	19 32 49 53 60 72 91 103 118 125 130-
			131 134 184 224 275 338 350 354 361-
			363 374 384 390 394 396 431-432 434-
			435 445 468 549 621 732 734-736 745
			760-761 764 768-769 775 787 806 811
			818 887 903 906 918 930 942 947 957
			973 977
- J-14 hi	Clantoch	ABR008	2-3 9-11 14 17 21 23-25 28-29 31-35 37
adult brain	Clontech	ADKUU	41-42 45 47-48 56-57 65-66 69-70 72 75
			77-78 88 91-92 97-99 101 103 112-115
		•	
			118-128 130-131 135 138-140 142 144-
			146 148 152 156-157 159-160 163 168
			172 174 176 178-180 182-190 194 196-
			198 200-201 204 209-214 218 220-225
			228-230 232-233 238-240 243-244 246
			254-256 260-264 270 272-274 278-279
			282-285 289-291 293-294 296-297 301
*			303-306 312-314 317 321-322 325-328
			334 336 338 340-342 344 346 348 350-
	ĺ		352 354 356-358 363 366 369-374 376
			379-381 383-386 388-394 398-399 402-
1	}	1	317-301 303-300 300-37+ 370-377 402*

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			403 405 409-412 414 418-421 423-424
			426-427 430 433-437 443 445-450 452
	•		456-457 460 462 464 471 479 482-483
			485 488 490-498 505 507 510 516 519-
			522 524 527-532 535 538-539 542-545
			548 551 553 555 561-562 566 569 571
			574 580-583 588-589 593 597 601-608
			611-612 614-615 617-618 621-622 624
			630-635 642 644 646-648 650-652 655
			657 659-661 664-665 668 672 674 689
			693-699 701-702 708 711 715 717 724
			728-730 732 734-735 738-740 745 747-
			750 753-755 757 761 763-764 766-769
			772-773 775 780-781 789-791 793-795
			799-800 802-806 809 812 818-819 821-
			822 826 829-830 832 834-835 841 843
			845 856 858-859 861 864 866 870 872
			876 880 883 885 887 893-898 902 906-
į			916 918 921 925-926 930-931 933 942-
			943 946 948 950-951 953-954 958-960
			962-965 967 969-970 972 977
adult brain	Clontech	ABR011	57 196 270 304 344 436 834
adult brain	BioChain	ABR012	14 82 121-122 168 691
adult brain	Invitrogen	ABR013	72 108 263 270 336 425 492-494 732 787
			790 826 880
adult brain	Invitrogen	ABR014	293 394 399 764 768-769 928 967
adult brain	Invitrogen	ABR015	738-739 764
adult brain	Invitrogen	ABR016	320 374 396 399 405 684 742-743 767
			931 947 967
adult brain	Invitrogen	ABT004	21 33-34 37-38 47 52 57-58 69 72 91-93
			109 119 122-124 126-127 135 142-143
			158 167-168 185-188 194 200 212 232
			242 246 255 258 270 277 279 293 301
			312-313 319 322-323 331 341 346 348
			371 374 388 391 394 399 401 409 411
			429 436-437 456 462 477 488 496 498
			510 512 515 539 542 545 549 559 563
			573 579 587 589 601-605 612 620-621
			624 640 643 647 681 715 723 728 732
			735-736 740 745 748 753 766 785-786
			792-793 797-801 812 822 829-831 853-
			856 859 876-877 884 893-894 908-909
			918 925 933 950 969 978
cultured	Strategene	ADP001	4 28-29 69 93 114 121 132-133 135 151-
preadipocytes ·			152 159 167 172 178 181 184 190 194-
		1	195 203-204 209 217 219 240 248 260-
			262 267 273-274 277 282 297 301 304
]			312 314 326-327 361-362 371 374 388
			394 401 403 405 411 420 437 453 466-
			467 470 474 478 496 507-509 517 530
			532-533 584 588 593 602-603 608 610
1		1	617-621 630-631 633 639 642-643 661

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 602 720 746 761 765 760 824 842 848

			693 729 746 761 765 769 834 842 848
			887 907 923 947-950 957 967 969
adrenal gland	Clontech	ADR002	1 3 12-13 21 23-24 27-29 67 74 78 103-
dustan B			105 108-109 113 115 118 120-121 128-
			133 149 156 160 172 177 182 214 217
			223 232-233 247 254 269-270 273-274
			277 283 285 288 298-299 308 317 319
			328 338 340 342 361-362 364 372 376-
			377 382 384 401-402 405-406 416 420
			431 437 444 446 448 457 462 484 500
			507 517 524 532-533 539 545 554 561-
			562 564 588 597 602-603 606-607 635
			642 646 649 658 664 674 693 703 730
			740 745 752 759 765 767 775 779 799
			809 817-818 839 845 856 859 863 887
			890-891 896 948 953 958 961-963 973
adult heart	GIBCO	AHR001	1 3-4 8 10 14 20-21 25 28-29 33-34 37-38
	GIBCO		41 48 54-57 65 69-72 75 78 80 82-83 97
	ł		99-100 108 112-115 117-121 123-124
			128-133 141 144-146 149 152 159 162-
			163 168 172 176 179 181 184 186-187
			190-191 201 203 208-209 212 216-218
			221 223 227 229 233 244 247 249 253-
			255 258 263-264 267 269-270 274 278
			280-282 285 289 291 295 297-299 301
			303-304 308 313 317 321-322 326 328
			334 344 348 352 358 361-363 370-371
			380 382-383 388 394-396 398 401 403
			405-406 410-416 423 425-427 430-431
			436 452-453 464-465 470-474 481-484
			487-488 490 492-494 496 499-500 505-
·			506 508-509 514 523 529-530 533 547-
			548 553 558 563-565 577-578 586-588
			590 593 597 601-603 606-608 610-613
			617-619 621-622 626-628 637-638 642-
			644 652 658 661 672 682-683 688 691
·			693 697 699 708 711 713 715 732 737
			745 747-748 750-753 759 761 765 768-
			770 775 790 802-803 814-815 818-819
1			830 837 839-840 842 845 848 859 861-
			862 867 876-877 887 891-892 896 900-
			901 903 905-906 908-909 919-920 922
			925 928 936 939-940 946-947 950 953
			959 967 970-971 973 977
adult kidney	GIBCO	AKD001	1.3 8 12-14 17 19-25 28-29 33-34 37-39
			41 46-48 50 52 55-60 62 65-67 69 71-72
			75 77-78 82 84 89-90 93 97 108-110 114-
			116 118-121 123-125 128 130-133 135
			138 144 146 149 156 159-161 163-164
†	Í		167-172 176 179 184 186-187 189-190
			194 196 200-202 204 209 211-212 216-
			217 219 221 223-224 229 232-235 244
L	<u> </u>	<u> </u>	41, 41, 441 LLJ-LLT LL, LJL-LJJ LTT

			247 250 253 255-256 258 263-264 268-
			272 274 277-281 283 286 288-290 292
			294-295 297 301 303-309 311-314 316
			319-323 325 328-338 342 348-349 352
	•		354-355 358 361-363 365 370-371 373
			376-378 380 382-383 388 395-399 401-
ļ			403 405-406 409-413 416 418-420 425-
			428 430-431 440 442 452-454 462 464-
		Δ.	465 470 472-474 477 479 481 483-485
1	-	**	487-489 492-495 498-500 504 506 510
			517 522 525 529-530 532-533 539 542-
			543 547 551-552 558 560-564 569-570
			573-574 577-578 580-583 585-590 594-
			596 601-608 610-613 617-621 624 626-
			628 630-631 634-636 639 642-643 648
			652 656 658 664-665 676-677 679 681
			688-691 693 697 699 708 711 715 717
1			720-722 724 729-732 738-741 747-748
'			751-753 761 765 770-778 780 784 789
	!		791 793 797 804 813 817 823-824 834
			837 839 842-843 845 848 859 861-862
			864 867 870 876-877 887 889 892-894
			896-897 900-901 903 907 913-915 918
			921 923 925 929-930 932 939 942 946-
İ			947 949-950 953 958-959 961-963 967
			1 969 972 977
adult kidney	Invitrogen	A K T 0 0 2	969 972 977
adult kidney	Invitrogen	AKT002	1 3 16 21 30 32 35 38-41 46-47 56 77 92
adult kidney	Invitrogen	AKT002	1 3 16 21 30 32 35 38-41 46-47 56 77 92 109 123-124 130-131 146 149 161 167-
adult kidney	Invitrogen	AKT002	1 3 16 21 30 32 35 38-41 46-47 56 77 92 109 123-124 130-131 146 149 161 167- 168 172 176 190 209 212 234-235 258
adult kidney	Invitrogen	AKT002	1 3 16 21 30 32 35 38-41 46-47 56 77 92 109 123-124 130-131 146 149 161 167- 168 172 176 190 209 212 234-235 258 279 292 301 303 308 314 333 355 363
adult kidney	Invitrogen	AKT002	1 3 16 21 30 32 35 38-41 46-47 56 77 92 109 123-124 130-131 146 149 161 167- 168 172 176 190 209 212 234-235 258 279 292 301 303 308 314 333 355 363 372 380 383 396 399 402 418-419 426-
adult kidney	Invitrogen	AKT002	1 3 16 21 30 32 35 38-41 46-47 56 77 92 109 123-124 130-131 146 149 161 167- 168 172 176 190 209 212 234-235 258 279 292 301 303 308 314 333 355 363 372 380 383 396 399 402 418-419 426- 427 431 448 454 461 471-474 488-489
adult kidney	Invitrogen	AKT002	1 3 16 21 30 32 35 38-41 46-47 56 77 92 109 123-124 130-131 146 149 161 167- 168 172 176 190 209 212 234-235 258 279 292 301 303 308 314 333 355 363 372 380 383 396 399 402 418-419 426- 427 431 448 454 461 471-474 488-489 495 498 504 506 508-509 520-521 530
adult kidney	Invitrogen	AKT002	1 3 16 21 30 32 35 38-41 46-47 56 77 92 109 123-124 130-131 146 149 161 167- 168 172 176 190 209 212 234-235 258 279 292 301 303 308 314 333 355 363 372 380 383 396 399 402 418-419 426- 427 431 448 454 461 471-474 488-489 495 498 504 506 508-509 520-521 530 537 539-541 545 547 563 582-583 592
adult kidney	Invitrogen	AKT002	1 3 16 21 30 32 35 38-41 46-47 56 77 92 109 123-124 130-131 146 149 161 167- 168 172 176 190 209 212 234-235 258 279 292 301 303 308 314 333 355 363 372 380 383 396 399 402 418-419 426- 427 431 448 454 461 471-474 488-489 495 498 504 506 508-509 520-521 530 537 539-541 545 547 563 582-583 592 613 617-618 621 623-624 633 655 688
adult kidney	Invitrogen	AKT002	1 3 16 21 30 32 35 38-41 46-47 56 77 92 109 123-124 130-131 146 149 161 167- 168 172 176 190 209 212 234-235 258 279 292 301 303 308 314 333 355 363 372 380 383 396 399 402 418-419 426- 427 431 448 454 461 471-474 488-489 495 498 504 506 508-509 520-521 530 537 539-541 545 547 563 582-583 592 613 617-618 621 623-624 633 655 688 690 693 699 704 713 732 745 752-753
adult kidney	Invitrogen	AKT002	1 3 16 21 30 32 35 38-41 46-47 56 77 92 109 123-124 130-131 146 149 161 167- 168 172 176 190 209 212 234-235 258 279 292 301 303 308 314 333 355 363 372 380 383 396 399 402 418-419 426- 427 431 448 454 461 471-474 488-489 495 498 504 506 508-509 520-521 530 537 539-541 545 547 563 582-583 592 613 617-618 621 623-624 633 655 688 690 693 699 704 713 732 745 752-753 761 766-768 770 784 789 797 837 842
adult kidney	Invitrogen	AKT002	1 3 16 21 30 32 35 38-41 46-47 56 77 92 109 123-124 130-131 146 149 161 167- 168 172 176 190 209 212 234-235 258 279 292 301 303 308 314 333 355 363 372 380 383 396 399 402 418-419 426- 427 431 448 454 461 471-474 488-489 495 498 504 506 508-509 520-521 530 537 539-541 545 547 563 582-583 592 613 617-618 621 623-624 633 655 688 690 693 699 704 713 732 745 752-753
adult kidney	Invitrogen	AKT002	1 3 16 21 30 32 35 38-41 46-47 56 77 92 109 123-124 130-131 146 149 161 167- 168 172 176 190 209 212 234-235 258 279 292 301 303 308 314 333 355 363 372 380 383 396 399 402 418-419 426- 427 431 448 454 461 471-474 488-489 495 498 504 506 508-509 520-521 530 537 539-541 545 547 563 582-583 592 613 617-618 621 623-624 633 655 688 690 693 699 704 713 732 745 752-753 761 766-768 770 784 789 797 837 842
adult kidney	Invitrogen	AKT002	1 3 16 21 30 32 35 38-41 46-47 56 77 92 109 123-124 130-131 146 149 161 167- 168 172 176 190 209 212 234-235 258 279 292 301 303 308 314 333 355 363 372 380 383 396 399 402 418-419 426- 427 431 448 454 461 471-474 488-489 495 498 504 506 508-509 520-521 530 537 539-541 545 547 563 582-583 592 613 617-618 621 623-624 633 655 688 690 693 699 704 713 732 745 752-753 761 766-768 770 784 789 797 837 842 848-849 866-867 877 887 893-894 903
	Invitrogen	AKT002	1 3 16 21 30 32 35 38-41 46-47 56 77 92 109 123-124 130-131 146 149 161 167- 168 172 176 190 209 212 234-235 258 279 292 301 303 308 314 333 355 363 372 380 383 396 399 402 418-419 426- 427 431 448 454 461 471-474 488-489 495 498 504 506 508-509 520-521 530 537 539-541 545 547 563 582-583 592 613 617-618 621 623-624 633 655 688 690 693 699 704 713 732 745 752-753 761 766-768 770 784 789 797 837 842 848-849 866-867 877 887 893-894 903 914-915 925 929-930 937 944-945 947-
adult kidney			1 3 16 21 30 32 35 38-41 46-47 56 77 92 109 123-124 130-131 146 149 161 167- 168 172 176 190 209 212 234-235 258 279 292 301 303 308 314 333 355 363 372 380 383 396 399 402 418-419 426- 427 431 448 454 461 471-474 488-489 495 498 504 506 508-509 520-521 530 537 539-541 545 547 563 582-583 592 613 617-618 621 623-624 633 655 688 690 693 699 704 713 732 745 752-753 761 766-768 770 784 789 797 837 842 848-849 866-867 877 887 893-894 903 914-915 925 929-930 937 944-945 947- 949 955 961 967 984
			1 3 16 21 30 32 35 38-41 46-47 56 77 92 109 123-124 130-131 146 149 161 167- 168 172 176 190 209 212 234-235 258 279 292 301 303 308 314 333 355 363 372 380 383 396 399 402 418-419 426- 427 431 448 454 461 471-474 488-489 495 498 504 506 508-509 520-521 530 537 539-541 545 547 563 582-583 592 613 617-618 621 623-624 633 655 688 690 693 699 704 713 732 745 752-753 761 766-768 770 784 789 797 837 842 848-849 866-867 877 887 893-894 903 914-915 925 929-930 937 944-945 947- 949 955 961 967 984 1 3 14 18 28-29 38 54-56 59 92 110 114-
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			1 3 16 21 30 32 35 38-41 46-47 56 77 92 109 123-124 130-131 146 149 161 167- 168 172 176 190 209 212 234-235 258 279 292 301 303 308 314 333 355 363 372 380 383 396 399 402 418-419 426- 427 431 448 454 461 471-474 488-489 495 498 504 506 508-509 520-521 530 537 539-541 545 547 563 582-583 592 613 617-618 621 623-624 633 655 688 690 693 699 704 713 732 745 752-753 761 766-768 770 784 789 797 837 842 848-849 866-867 877 887 893-894 903 914-915 925 929-930 937 944-945 947- 949 955 961 967 984 1 3 14 18 28-29 38 54-56 59 92 110 114- 115 130-131 146 149 156 159 164 167 176 184 209 217 234-236 240 255-256 258 263-264 269 271 276 280-281 297 305 308 312 314 322 325 332 336 344 353 361-362 388 401 410 420-421 426- 427 431 465 469 474 484 498 500 506 508-509 517 530 532 573 592 596 613 619-620 623 626-628 638 658 679 681
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			967
lymph node	Clontech	ALN001	3 10 110 146 160 168 196 209 221 269
			278 301 336 348 394 405 411 420 422
			459 464 474 485 503 506-507 532 563
			582 619 623 630-631 642 669 684 697
			713 715 727 747 767 769 789 825 839
			842 849 887 896 913 921 925
young liver	GIBCO	ALV001	3 14 16 37-38 41 51 56 60 97 104-105
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	}		149 152 169-172 176 184 189-190 200
			209 212 216 218 228 232 255 258 263
			270-271 275 285-286 292 295 298-299
			301 304 314 341 358 365 368 376 400
			410-412 431 474 481-482 485 496 500
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	1		504-505 517 520-522 524 530 532-533
			547 551 563 581 583 610-611 621 624
		1	635 643 691 708 711 715 720 752 755
			761 768 796-797 811 818 830 845-847
<u> </u>			852 864-865 867-869 896 899 910-911
			949 958 965 969 972-973
adult liver	Invitrogen	ALV002	3 37 42 56 60 71 82 104-105 114-115
			117-118 125 130-131 134-135 164 169-
			172 176 179 200 203-204 212 217 223
		•	226 232 237 244 263 274-275 292 301
			310-312 314 317 349 354 364 368 372
			376 398-399 402 426-427 439 442 451
			458 465 474 482 485 490 506 515 525
			527 545 547 552 568 571 573-575 582
			587 594-595 604-605 608 610 621 630-
			631 634-635 637 657 664 690 693 699
			723 726 745 751 763 767 784 793 811
		1	822 845 848 852 856 861-862 864 892
			899 908-909 925 950 958 967 983
		1 7 7 7 0 0 0	
adult liver	Clontech	ALV003	60 134 169-171 275
adult ovary	Invitrogen	AOV001	1 3 9-10 12-14 16 18 20 22-25 28-29 33-
			35 37 39 41-42 46 48-50 55-57 59 63-67
			69 71-72 75 77-80 82 88-89 92 101 103-
		ļ	106 108-110 113 115 119-121 123-126
			128-133 135 138 142-146 149 151-152
·			159-161 167-168 172 174 176-177 179
			181 184-190 194 198 200 203 208-209
			211-212 214 217 219 221 224 226 232-
			235 240-242 246-247 249 251 254-255
			258-259 264 269-271 274 276-277 279-
		*	283 285 288 290 293-294 297 301-304
1			306-308 311 314 319-322 325-326 328-
			329 331-332 335-338 341-342 344 348
			354-358 361-363 365 368 370-372 374
			376 379-380 382-383 388 394-396 398-
			399 401-402 405-406 409-412 416 418-
			421 423 425-433 438 442-443 449-452
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			482-484 488 490 492-496 498 500-504
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			532-533 537 539-542 545 551 555 558
			560-565 569 571 573 577-578 581-583
			585-590 592-593 596-597 600-605 608
			610-611 613-614 617-628 633-637 639
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			664 668-670 672 674 679 681 684 688
			691 693 697-699 701-702 713 717 721-
i			722 724 729-732 738-744 747-750 752-
			753 755 759 761 765 767-774 779-780
	1		783-784 789 793 795-797 801 813-818
			823-824 828 830-832 834 837 839 841-
			842 845 848-851 856 859 862 864 866-
			867 870-871 874-878 881-883 887-889
			891 893-894 896-897 901 903 906-911
			913 919-922 925 928 930 936 939-940
	1		943-944 946-947 949-950 952-953 955
			957-958 962-963 965 967 969 971 973
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adult placenta	Invitrogen	APL001	41 56 67 253 301 304 334 380 383 451
			474 479 500 577-578 643 648 729 767
			856 859 866 873 962-963
placenta	Invitrogen	APL002	3 21 31 38 63-64 78 135 143 168 186-187
			212 232 244 263 280-281 334 336 344
			348 371 374 394 399 461 490 582 588
			602-607 610 620 699 745 769 793 817
			822 859 897-898 923 928 931 943 949
			969 973
adult spleen	GIBCO	ASP001	1 3 21-22 46 52 54-55 57-58 61-62 72 74
			78 82 88 118 121 130-131 137 152 159
			168 172 189 203 209 217 223 234-235
			252 255 263 269 271 274 282 288 290
	İ		301 314 322 335 350 363 394 403 405-
			406 410-412 415 431 459 464 472-474
			482 488 500 506 510 514 517 532 537
			542 561-563 589 593 602-603 610 613
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			676 679 681-682 684 689 691-692 697
			699 715 720 723 729 747-748 769-770
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testis	GIBCO	ATS001	6 22 28-29 33-34 41 48 52 62 65 72 97
			106 109 118 132-133 145-146 168 172
1			176 183 185 189-191 195 209 211-212
			214 221 223 230 254-255 258 263 269
			283 297 312 314 321 342 352 361-362
			365 380 383 388 395 401 405-406 412
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Genomic DNA from BAC 63I18	Research Genetics (CITB BAC Library)	BAC001	515
Genomic DNA from BAC 393I6	Research Genetics (CITB BAC Library)	BAC002	640
Genomic DNA from BAC 393I6	Research Genetics (CITB BAC Library)	BAC003	640
adult bladder	Invitrogen	BLD001	50 55 66 71 111 143-144 148 160 201 209 223 255-256 280-281 286 305 315 319 340 394 431 442 488 497 505 518 552 588-589 621 636 664 676 715 738-739 769 790 824 837 845 877 887 936 940 948 962-963 967
bone marrow	Clontech	BMD001	3 10-13 16 18 20-21 25 28-29 31-34 41 45 48 52 54-55 57 59 61 65 67 72-73 75 78 80 82 84 99 103 108 110 114-115 118-120 123-124 128 130-133 143-144 148 152 159-161 163 168 172 174 176 178 190 192 198 203 209 211 217-218 221 223-224 227 233-236 244 247 249 252 254 258 260-262 267 269 272 278 280-281 284-285 288 290 294-297 301 304 308 314 317-318 320-321 325 328-330 333-335 349 351-354 358 363 365 367 377 382 388 394-397 400 405 408 410-412 418-421 425-428 431 433 435 442 449-450 453 455 459 464 468-470 474 478-479 481 484 490 496 504 506 508-509 511 519-521 530 532 539 553 558-559 561-563 580 582 586 592 599 608 610 613-614 617-619 623 625-628 635 638 641-643 658 664 672 682 699 711 713 717 731 734 740 742-743 745 761 768-771 774 776-778 784 787 789 813 817-818 822 834 839-840 842 848 862 866 870 876 885-887 891 896-898 900 903 906 913 919 921-922 927-928 939 944 947 950 953 959 961-963 967-968 970 973 977
bone marrow	Clontech	BMD002	3 9-10 15-19 30 33-34 39 45 54 57 63-64 71 82 102 116 119 130-133 148 152 156

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			159-160 168 176 182 224 254-255 271-
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1		1	367 370 372 387 394-395 399 403 405
			409 411 449-450 459 461 468 474 488-
			489 524 530 532 580-582 592 602-603
			611 617-618 621-622 630-632 642 661
			663 694 717 730 734 740 745 752 755
			761 767 769-771 775-778 784 787 811
			813 818 832 840 842 849 859 878 887
			893-894 896-898 903 906 908-909 923
		1	928 944 946-949 953 958-963 965 982
1	Olamas al	DM 0004	
bone marrow	Clontech	BMD004	54
bone marrow	Clontech	BMD007	766 887 928
adult colon	Invitrogen	CLN001	22 37 67 97 117 121 148-149 168 172 190
			200 204-205 232 244 263 268 292 301-
			302 363 377 384 452 455 459 470 530
			582 602-603 619 687 723 728 751 761
			831 861 887 914-916 934 955 969 984
Mixture of 16	Various	CTL016	358 740 760
tissues –	Vendors*		
mRNAs*	, 533330		
Mixture of 16	Various	CTL021	468 527 928
tissues -	Vendors*	012021	100 327 320
mRNAs*	Vendors		
adult cervix	BioChain	CVX001	1 3 10 14 22 28-30 37 41 47-48 51-52 54-
adult cervix	BioChain	CAYOU	
			57 71 82 89-90 92 106 108 110-111 117-
	l	}	118 121 129-131 135 141 143-146 160-
			161 164 168 172 177 189-190 193 195
			200 204 209 211-212 217 226 229-230
			232 234-235 240-242 246 254 260-263
			268-270 274 277 282 285 292 295 297
			305-308 314-316 319 328 343-344 348
			354 358 363 368 380 382-384 389 394
	[396 399 401 405-407 410 416 418-421
			428 430-431 437 442 453-454 459 464
			469 471-473 476 480 484 492-495 500
			504 506-509 516-517 526 530 532 545
			550-551 563-565 569 577-578 585-586
			590 608 611 613 619 621 623 628 630-
	}		631 634-637 641 643 648 656-658 664-
			665 674 679 682 689-690 693 700 703
		•	708 713 721-722 724 728 732 742-743
4	1		747 750 752 755 757 761 763 767-769

The 16 tissue-mRNAs and their vendor source, are as follows: 1) Normal adult brain mRNA (Invitrogen), 2) normal adult kidney mRNA (Invitrogen), 3) normal adult liver mRNA (Invitrogen), 4) normal fetal brain mRNA (Invitrogen), 5) normal fetal kidney mRNA (Invitrogen), 6) normal fetal liver mRNA (Invitrogen), 7) normal fetal skin mRNA (Invitrogen), 8) human adrenal gland mRNA (Clontech), 9) human bone marrow mRNA (Clontech), 10) human leukemia lymphablastic mRNA (Clontech), 11) human thymus mRNA (Clontech), 12) human lymph node mRNA (Clontech), 13) human spinal cord mRNA (Clontech), 14) human thyroid mRNA (Clontech), 15) human esophagus mRNA (BioChain), 16) human conceptional umbilical cord mRNA (BioChain).

			779-780 784 788 810-811 813-815 822
			834 836-837 839 848 861 866-867 871
			874 877 887 891-894 897-898 901 913
			916 919 921-922 925 946-947 953 958-
			959 967 969 973
diaphragm	BioChain	DIA002	3 39 184 203 431 563 848 967
endothelial	Strategene	EDT001	3 6 8-10 14 19-24 28-29 33-34 37 39 41
cells	S	3303	46 48 52 55-58 62-65 67 69 71-72 75 78
			80 82-83 87 101-102 108-109 114-115
			117 123-124 128 130-133 135 138 143
			145-146 149 156 159-160 167-168 172
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			194-195 200 203 208-209 212 216-217
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fetal kidney	Clontech	FKD001	3 31 33-34 38 48 54 72 160 208-209 211
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fetal kidney	Clontech	FKD002	19 474 726 903
fetal kidney	Invitrogen	FKD007	3 118 186-187 230 244 271 432 887 969
fetal lung	Clontech	FLG001	69 132-133 156 168 208-209 217 267 269
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	·	1	437 466-467 471 484 500 517 520-521
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			612 624 643 711 731 733-734 757 761
			769 774-775 794 824 864 904 906 910-
Į.		·	911 913 948 953 959 976 984
skeletal muscle	Clontech	SKM001	15 75 135 146 172 190 218 267 282 308
			410 426-427 474 505 588 620 623 658
			692 713 737 779 790 862 874 878 887
			952 962-963
skeletal muscle	Clontech	SKMs04	215
spinal cord	Clontech	SPC001	14 20-21 25 28-29 31 39 46 48 59 78 83-
spinar cord	Cionicon	51 0001	84 91-92 103 112-113 135 160 168 172
			176 188 190 205 209 229 232 258 285
			301 308 312-314 321 323 329 346 374
			377 380 383 388 394 398 406 409-410
1		1	431 449-450 453 455 466-467 470-471
			484-486 488 495 497 500 503 508-509
			524 537 539 558 581 586 604-605 611
			619 623 630-631 633 656 663 711 715
			729 736 740-741 761 767 769 776-778
	1		723 100 110 110 110 110 110 110
			780 818 822 831 835-836 840 843 859
			861 871 875 887-888 897 906-907 913
			919-920 928 931 953 958
adult spleen	Clontech	SPLc01	3 6 12-13 66 130-131 178 365 403 431
			461 558 610 715 797 809 876 947 967
stomach	Clontech	STO001	35 114 130-131 144 155 176 189 206-207
			249 260-262 336 382 398 425 431 453
			461 483 496 500 527 530 580 642 657
			663 669 748 765 768 802-803 839 891
			942 981
thalamus	Clontech	THA002	30-32 48 66 109 127 130-131 135 142
			145 156-158 168 172 174 185 199 224-
}			225 233 246 277 282 286 293 322 332
			334 346 374 384 400 402 420 424 435-
			437 446 466-467 485 503 506 527 542
			549 572 612 615 622 624 633 643-644
			658 676 736 790 794 824 831 835 896
		,	907 950 969
thymus	Clonetech	THM001	10 16 20 28-29 32 37 41 52 57 66-67 74-
anymus .			75 110 118 121 129-131 141 151 159-160
			208 211 218 247 269 289 295 297 320
			325 354 358 365 367 372 378 388-389
			395 398 411-412 420 423 435 452 500
			508-509 517 524 532 537 551 558 560
	<u> </u>		JUU-JUJ J11 J27 JJ2 JJ1 JJ1 JJU JUU

			569 577-578 582 586 598 608 611 622
			643 684 715 721-723 728 740 766 772-
	į		773 795 834 837 849 864 885 900 921
			946 948 958 962-963 965 972-973 982
thymus	Clontech	THMc02	1 3 9-11 16 21 27 32-34 38-39 51 55-57
			66 72 74 77-78 80 82 89-90 101 112 115
			118-119 121 123-124 126 138 144 152
	[159 168 174 176 178 186-1-8 197 200
		-	208 212-214 217 225 233 243-244 246
			254 256-262 279 282 285 288-289 296-
		'	297 313-314 322 334 343 354-355 358-
			359 363-364 367-368 372-373 382 387-
•			389 395 400 402 411 414 426-427 437
			440 442 449-450 454 457 462 464 469
			474 479 481 485 490-491 506 508-509
			511 517 522 526 528 532 542 551 554
			561-562 564 566-570 580-582 585 589
	İ		597 599-600 602-608 611 613-614 619-
			621 625 628 630-631 644 646 655 669
			672 677 684 686-693 697 713 717 720
			728 740 746 749 760-762 767 771 775
	İ		794 797 804 808 811 816 818-819 837
			840 859 880 883 887-888 896-897 903
			908-911 913 916 924 936 947-948 950
			962-963 965 967 970
thyroid gland	Clontech	THR001	3 8-9 14-15 19-22 28-29 39 41 55-56 66
,	}		69 71-72 78-79 97 104-105 109 113 115
			119 121 123-124 130-133 135 138 143-
			144 146 148 151-152 156 159-163 165
			168 172 174 177 183-184 196 199-200
			203 209 211 215-218 228-229 232-236
			244 254-255 258 273 282 290 292 294
	•		297 303-306 308 311 317-318 322-323
			325-326 334-335 340 342 348 354 358
•			373 377 381-382 387 394 398 401-402
			405-406 409-412 416 422 425-427 429-
			431 440 449-453 462 466-468 474 478-
			479 481-484 490 492-496 500-501 505-
			506 517-518 522-525 532 537 540-541
			545 551 558 560 563-564 580 583 587-
			589 593 597 599 606-607 610 617-621
			625-628 633 635 641-643 658-659 664-
			669 674 682 686 688-691 696 699 715
			724 730 740 742-743 747 750 752 759
			761 765-766 768-769 779 789 796 802-
			803 813 818-819 822 831 837 843 845
			848-849 862 864 868-869 871 874 876-
			877 887 893-894 896-897 907-909 912
			919-921 923 925 928 936 940-942 944
			946-947 950 953 955 958-959 962-963
	[967 969 973 981
trachea	Clontech	TRC001	33-34 55-56 69 74 163 172 190 209 212
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267 270 297 305 314 352 413 426-427 466-467 500 502 504 580 586 610 613 633 642 688 691 711 724 738-739 774 782 816 820 839 848 862 868-869 914-915 928 968 Clontech UTR001 4 9 18 37 63-64 74 108 114-115 130-131 uterus 160 166 179 184 190 209 233 249 269 285 301 314 327 337 348 384 394 399-400 403 406 411 425 431 434 437 440 462 474 485 490 508-509 526 532 579 617-619 636 642-643 672 761 769 793 837 849 864 887 903 906 928 934 947 967

TABLE 2

SEQ ID NO:	ACCESSION NUMBER	SPECIES	DESCRIPTION	SMITH- WATERMAN SCORE	% IDENTITY
1	L06175	Homo sapiens	occurs in MHC class I region; ORF	308	98
2	Y70775	Homo sapiens	Follistatin-related protein zfsta.	3094	98
3	X15187	Homo sapiens	precursor polypeptide (AA -21 to 782)	4112	100
4	AF110640	Homo sapiens	orphan seven-transmembrane receptor	344	100
5	G03798	Homo sapiens	Human secreted protein, SEQ ID NO: 7879.	158	72
6	W85607	Homo sapiens	Secreted protein clone da228_6.	1477	100
7	Y30162	Homo sapiens	Human dorsal root receptor 4 hDRR4.	884	88
8	Y15227	Homo sapiens	Leu1	391	100
9	Y28817	Homo sapiens	pt326_4 secreted protein.	3338	100
10	X92106	Homo sapiens	bleomycin hydrolase	2445	100
11	Y15228	Homo sapiens	Leu2	445	100
12	U27838	Mus musculus	glycosyl-phosphatidyl-inositol- anchored protein homolog	432	34
13	U27838	Mus musculus	glycosyl-phosphatidyl-inositol- anchored protein homolog	320	27
14	Y71062	Homo sapiens	Human membrane transport protein, MTRP-7.	2323	99
15	U96781	Homo sapiens	Ca2+ ATPase of fast-twitch skeletal muscle sacroplasmic reticulum, adult isoform	5145	100
16	M16653	Homo sapiens	pancreatic elastase IIB zymogen	1435	99
17	Y13398	Homo sapiens	Amino acid sequence of protein PRO346.	1749	99
18	Y02283	Homo sapiens	Secreted protein clone br342_11 polypeptide sequence.	1399	99
19	Y53030	Homo sapiens	Human secreted protein clone d24_1 protein sequence SEQ ID NO:66.	1371	100
20	AL031320	Homo sapiens	dJ20N2.5 (novel protein similar to fucosidase, alpha-L-1, tissue (EC 3.2.1.51, alpha-1-fucosidase fucohydrolase))	2597	99
21	B01384	Homo sapiens	Neuron-associated protein.	1876	100
22	Y68778	Homo sapiens	Amino acid sequence of a human phosphorylation effector PHSP-10.	2470	100

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SEQ ID NO:	ACCESSION NUMBER	SPECIES	DESCRIPTION	SMITH- WATERMAN SCORE	% IDENTITY
23	Y55935	Homo sapiens	Human KHS2 protein.	4781	99
24	Y55935	Homo sapiens	Human KHS2 protein.	2807	100
25	AC024792	Caenorhabditis elegans	contains similarity to TR:O95029	463	31
26	Y07972	787	Human secreted protein fragment	1540	100
27	X97630	Homo sapiens	serine/threonine protein kinase	3781	98
28	AF150755	Mus musculus	microtubule-actin crosslinking factor	3514	68
29	AF150755	Mus musculus	microtubule-actin crosslinking factor	3725	70
30	Z38011	Mus musculus	DMR-N9	2988	86
31	AJ000522	Homo sapiens	axonemal dynein heavy chain	6058	99
32	AF037256	Mus musculus	ES2 protein	2260	91
33	S62140	Homo sapiens	TLS=nuclear RNA-binding protein	2917	100
34	S62140	Homo sapiens	TLS=nuclear RNA-binding protein	2890	98
36	AB038237	Homo sapiens	G protein-coupled receptor C5L2	1767	100
37	D79994	Homo sapiens	similar to ankyrin of Chromatium vinosum.	6089	99
38	X63380	Homo sapiens	serum response factor-related protein	1966	99
39	AL022072	Schizosacchar	lipoic acid synthetase	1067	61
	¥20000	omyces pombe		0751	100
40	J03930	Homo sapiens	alkaline phosphatase	2751 1088	100 98
41	AF132968	Homo sapiens	CGI-34 protein hypothetical protein	2208	100
42	AL117637 AL021393	Homo sapiens Homo sapiens	bK747E2.1 (novel protein)	1526	100
43	X68011	Homo sapiens	ZNF81	1886	100
45	AC002464	Homo sapiens	organic cation transporter; 50%	2423	100
			similarity to JC4884 (PID:g2143892)		
46	W78245	Homo sapiens	Fragment of human secreted protein encoded by gene 19.	1949	100
47	Y41765	Homo sapiens	Human PRO1083 protein sequence.	3604	100
48	AF097330	Homo sapiens	H1 chloride channel; p64H1; CLIC4	1305	99
50	U09413	Homo sapiens	zinc finger protein ZNF135	1361	57
51	AF061812	Homo sapiens	keratin 16	2374	100
52	W63681	Homo sapiens	Human secreted protein 1.	1326	99
53	AB035303	Homo sapiens	cadherin-10	4094	100
54	A12022	synthetic construct	MRP-8	485	100
55	AL121897	Homo sapiens	bA392M18.3 (KIAA0180)	1867	100
56	¥73330	Homo sapiens	HTRM clone 397663 protein sequence.	818	96
57	AF151018	Homo sapiens	HSPC184	955	100
58	AF125042	Homo sapiens	bisphosphate 3'-nucleotidase	1586	100
59	AF118670	Homo sapiens	orphan G protein-coupled receptor	1971	100
60	X04494	Homo sapiens	precursor polypeptide	1903	100
61	AF208865	Homo sapiens	EDRF	528	100
62	D15057	Homo sapiens	DAD-1	567	100
63	AF260665	Homo sapiens	histone acetyltransferase	1510	100
64	AF260665	Homo sapiens	histone acetyltransferase	1429	100
65 66	AJ277145 Y94950	Homo sapiens Homo sapiens	ras-related small GTPase RAB18 Human secreted protein clone dh1073_12 protein sequence SEQ ID NO:106.	1073 348	100
67	Y82744	Homo sapiens	DNA replication and repair associated protein (DRASP).	1028	100
68	Y44486	Homo sapiens	Human GPRW receptor polypeptide.	1721	100
69	AL031228	Homo sapiens	dJ1033B10.2 (WD40 protein BING4 (similar to S. cerevisiae YER082C, M. sexta MNG10 and C. elegans F28D1.1)	3196	100
	L	1			

SEQ ID NO:	ACCESSION NUMBER	SPECIES	DESCRIPTION	SMITH- WATERMAN SCORE	% IDENTITY
70	AJ276316	Homo sapiens	zinc finger protein 304	1751	52
71	Y18314	Homo sapiens	paraplegin-like protein	4146	99
72	AF157028	Homo sapiens	protein phosphatase methylesterase-1	2017	100
74	Y71082	Homo sapiens	Human B-aggressive lymphoma (BAL) protein.	1765	99
75	AF225420	Homo sapiens	AD025	734	100
76	X95235	Homo sapiens	transcription factor AP2	217	100
77	AF108420	Takifugu rubripes	1-aminocyclopropane-carboxilate synthase	733	56
78	G01349	Homo sapiens	Human secreted protein, SEQ ID NO: 5430.	650	99
79	AL117635	Homo sapiens	hypothetical protein	922	99
81	Z85986	Homo sapiens	dJ108K11.3 (similar to yeast suppressor protein SRP40)	865	77
82	AF183414	Homo sapiens	hemin-sensitive initiation factor 2a kinase	3231	99
83	G01143	Homo sapiens	Human secreted protein, SEQ ID NO: 5224.	495	98
84	U03985	Homo sapiens	N-ethylmaleimide-sensitive factor	3744	99
85	Y17791	Homo sapiens	VAX2 protein	1496	100
87	AF263538	Homo sapiens	growth differentiation factor 3	1944	99
88	Y19757	Homo sapiens	SEQ ID NO 475 from WO9922243.	1361	100
89	AF161493	Homo sapiens	HSPC144	1185	100
90	AF161493	Homo sapiens	HSPC144	856	100
91	B25780	787	Human secreted protein SEQ ID	647	41
92	U57344	Mus musculus	Meis3	1007	89
93	AF172854	Homo sapiens	cardiotrophin-like cytokine CLC	1197	98
94	AL390114	Leishmania	extremely cysteine/valine rich	223	29
		major	protein		
95	AB016886	Arabidopsis thaliana	contains similarity to adenylate kinase~gene_id:MCA23.18	287	38
96	AC005525	Homo sapiens	F22162_1	1855	96
97	B20997	Homo sapiens	Human nucleic acid-binding protein, NuABP-1.	3836	99
98	AJ006692	Homo sapiens	ultra high sulfer keratin	507	70
99	AF172264	Homo sapiens	Traf2 and NCK interacting kinase, splice variant 1	6942	99
100	L11239	Homo sapiens	homeobox protein	717	100
101	AC004890	Homo sapiens	similar to zinc finger proteins; similar to AAC01956 (PID:g2843171)	2154	98
102	AC003682	Homo sapiens	R28830_2	1287	48
103	AF201839	Rattus norvegicus	dynamin IIIbb isoform	4270	95
104	Y79510	Homo sapiens	Human carbohydrate-associated protein CRBAP-6.	1394	100
105	Y79510	Homo sapiens	Human carbohydrate-associated protein CRBAP-6.	1209	90
106	AL096748	Homo sapiens	hypothetical protein	1216	100
108	X97260	Homo sapiens	Metallothionein 2	381	100
109	AL034422	Homo sapiens	dJ1141E15.2 (novel protein)	433	100
110	AF191338	Homo sapiens	anaphase-promoting complex subunit 4	683	100
111	AL021712	Arabidopsis thaliana	putative protein	185	26
112	AF250138	Homo sapiens	small stress protein-like protein HSP22	1063	100
113	AL109976	Homo sapiens	dJ794I6.1.1 (novel protein)	4176	99
114	Y36151	787	Human secreted protein	668	100

SEO	ACCESSION	SPECIES	DESCRIPTION	SMITH-	
ID NO:	NUMBER		DESCRIPTION	WATERMAN SCORE	identity
115	AF110399	Homo sapiens	elongation factor Ts	1666	100
116	AF210317	Homo sapiens	facilitative glucose transporter family member GLUT9	2052	99
117	Y73328	Homo sapiens	HTRM clone 082843 protein sequence.	931	100
118	X04085	Homo sapiens	catalase	2846	100
119	AF147717	Homo sapiens	ubiquitin C-terminal hydrolase UCH37	1695	100
120	X73882	Homo sapiens	microtubule associated protein	3801	99
121	AC004882	Homo sapiens	similar to CAA16821 (PID:g3255952)	3223	100
122	M93311	Homo sapiens	metallothionein-III	421	100
123	G03827	Homo sapiens	Human secreted protein, SEQ ID NO: 7908.	557	94
124	G03827	Homo sapiens	Human secreted protein, SEQ ID NO: 7908.	222	53
125	AF232009	Homo sapiens	peroxisomal trans 2-enoyl CoA reductase	1565	99
126	AB004906	Ipomoea purpurea	transposase	146	20
127	M60165	Homo sapiens	guanine nucleotide-binding regulatory protein 2	1832	99
128	Y10319	Homo sapiens	carnitine carrier	1592	100
129	U75467	Drosophila melanogaster	Atu	937	36
130	Z21507	Homo sapiens	human elongation factor-1-delta	494	87
131	Z21507	Homo sapiens	human elongation factor-1-delta	938	100
132	Y58633	Homo sapiens	Protein regulating gene expression PRGE-26.	6745	100
133	Y58633	Homo sapiens	Protein regulating gene expression PRGE-26.	4818	95
134	M13692	Homo sapiens	alpha-1 acid glycoprotein precursor	1064	99
135	U72970	Sus scrofa	calcium/calmodulin-dependent protein kinase II isoform gamma-B	2723	99
136	G03213	Homo sapiens	Human secreted protein, SEQ ID NO: 7294.	450	100
137	AC005102	Homo sapiens	small inducible cytokine subfamily A member 24	627	99
138	AF155648	Homo sapiens	putative zinc finger protein	5855	92
139	AF144638	Homo sapiens	sphingosine-1-phosphate lyase	2977	100
140	AF152318	Homo sapiens	protocadherin gamma A1	4778	100
141	B08517	Homo sapiens	Amino acid sequence of a beta- tubulin antigen.	5841	100
142	X56667	Homo sapiens	calretinin	1410	99
143	X92763	Homo sapiens	tafazzins	1605	100
	Y95293	Homo sapiens	Human GEF containing NEK-like kinase substrate sGNK.	4092	99
145	AF226046	Homo sapiens	GK003	1198	100
146	M22877 AJ272212	Homo sapiens	cytochrome c	554	98
147	AB026491	Homo sapiens Homo sapiens	protein serine kinase PICK1	2196	100
149	AB020491 AB018580	Homo sapiens	hluPGFS	2114 1699	98 100
150	X91868	Homo sapiens	six1	1509	100
151	AF266505	Mus musculus	pseudouridine synthase 3	2135	84
152	U29170	Drosophila	ANON-23D	883	43
153	G04075	melanogaster Homo sapiens	Human secreted protein, SEQ ID	567	99
154	AY009128	Homo sapiens	NO: 8156.	138	100
454	A 1 007 120	Tionio sapiens	10002	130	100

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SEQ ID NO:	ACCESSION NUMBER	SPECIES	DESCRIPTION	SMITH- WATERMAN SCORE	MENTITY
155	AF141315	Homo sapiens	alpha-1,4-N- acetylglucosaminyltransferase	1842	100
156	AF110645	Homo sapiens	candidate tumor suppressor p33 ING1 homolog	1294	99
157	AF159297	Zea mays	extensin-like protein	238	25
158	AL133325	Homo sapiens	dJ984P4.3 (Homeobox protein NKX2B)	1437	100
159	AF073298	Homo sapiens	small EDRK-rich factor 2	294	100
160	AC004858	Homo sapiens	U1 small ribonucleoprotein 1SNRP homolog; match to PID:g4050087	4032	100
161	AB012109	Homo sapiens	APC10	990	100
162	AL162751	Arabidopsis thaliana	putative protein	194	32
163	AJ005698	Homo sapiens	poly(A)-specific ribonuclease	3351	100
164	AF117646	Homo sapiens	long CBL-3 protein	2547	99
165	AC004002	Homo sapiens	similar to ciliary dynein beta heavy chain; 78% Similarity to P23098 (PID:g118965)	5065	100
166	M10942	Homo sapiens	human metallothionein-Ie	381	100
167	AF126484	Homo sapiens	CARD4	4961	100
168	AF161518	Homo sapiens	HSPC169	1604	100
169	M64983	Homo sapiens	fibrinogen beta chain	2482	100
170	M64983	Homo sapiens	fibrinogen beta chain	2679	100
171	M58514	Gallus gallus	fibrinogen beta chain	1059	78
172	AF078845	Homo sapiens	16.7Kd protein	786	100
173	AC004774	Homo sapiens	Dlx-6	923	100
174	Z98974	Schizosacchar omyces pombe	putative vacuolar protein sorting- associated protein	185	31
175	X56203	Plasmodium falciparum	liver stage antigen	283	23
176	W74726	Homo sapiens	Human secreted protein fg949_3.	1879	100
177	AJ222967	Homo sapiens	cystinosin	1920	100
178	AC024796	Caenorhabditis elegans	contains similarity to TR:076167	221	27
179	Y66632	Homo sapiens	Membrane-bound protein PRO276.	1370	100
180	AF151803	Homo sapiens	CGI-45 protein	215	28
181	G02694	Homo sapiens	Human secreted protein, SEQ ID NO: 6775.	283	100
182	Y17292	Homo sapiens	Human cell death preventing kinase (DPK-1) protein sequence.	2676	100
183	AF234765	Rattus norvegicus	serine-arginine-rich splicing regulatory protein SRRP86	148	27
184	AF151855	Homo sapiens	CGI-97 protein	1214	96
185	AF289664	Mus musculus	CYLN2	4673	90
186	AL022238	Homo sapiens	dJ1042K10.2 (supported by GENSCAN, FGENES and GENEWISE)	4059	100
187	AL022238	Homo sapiens	dJ1042K10.2 (supported by GENSCAN, FGENES and GENEWISE)	2332	100
188	X83543	Homo sapiens	APXL	8513	99
189	AF059569	Homo sapiens	actin binding protein MAYVEN	3106	99
190	M18135	Rattus norvegicus	smooth-muscle alpha tropomyosin	1306	95
191	AF242194	Drosophila melanogaster	brakeless-B	147	52
192	D30689	Bacillus subtilis	subunit of nitrite reductase	113	29
193	Y44984	Homo sapiens	Human epidermal protein-1.	538	97

SEQ ID NO:	ACCESSION NUMBER	SPECIES	DESCRIPTION	SMITH- WATERMAN SCORE	% IDENTITY
194	B25679	Homo sapiens	Human secreted protein sequence encoded by gene 15 SEQ ID NO:68.	760	100
195	AB020315	787	homologue of mouse dkk-1 gene:Acc	1466	100
196	U35730	Mus musculus	jerky	2021	75
197	AL136450	Homo sapiens	dJ510O21.1 (novel protein)	632	· 100
198	X56203	Plasmodium falciparum	liver stage antigen	512	24
199	Y70775	Homo sapiens	Follistatin-related protein zfsta.	2027	63
200	X87237	Homo sapiens	a-glucosidase I	4447	99
201	AF101078	Caenorhabditis elegans	CLU-1	1393	46
202	X04571	Homo sapiens	precursor polypeptide (AA -22 to 1185)	6611	100
203	X00474	Homo sapiens	pS2 precursor	466	100
204	AB029333	Halocynthia roretzi	HrPET-1	974	54
205	AF146019	Homo sapiens	hepatocellular carcinoma antigen gene 520	998	100
206	AF071002	Homo sapiens	minK-related peptide 1; MiRP1	632	100
207	AB038162	Homo sapiens	trefoil factor 2	744	100
208	U30521	Homo sapiens	P311 HUM	363	100
209	AB000911	Sus scrofa	ribosomal protein	782	100
210	AB021227	Homo sapiens	membrane-type-5 matrix metalloproteinase	3545	100
211	AF180920	Homo sapiens	cyclin L ania-6a	2722	100
212	AF105365	Homo sapiens	K-Cl cotransporter KCC4	5624	100
213	U29244	Caenorhabditis elegans	similar to human (TRE) transforming protein (PIR:S22157)	602	32
214	AL033538	Homo sapiens	dJ477H23.1 (novel protein)	3195	100
215	X52011	Homo sapiens	muscle determination factor	1262	100
216	AF083248	Homo sapiens	ribosomal protein L26 homolog	739	100
217	AF006751	Homo sapiens	ES/130	4793 3559	99
218	AB007859	Homo sapiens	KIAA0399 protein	826	100
219	AK026291	Homo sapiens	unnamed protein product	5851	97
221	Y84045	Homo sapiens	Splice variant of cancer associated polypeptide CH1-9a11-2.		100
222	Z67996	Homo sapiens	tenascin-R (restrictin)	7186. 846	100
223	AF134802	Homo sapiens	cofilin isoform 1	1611	99
224	Y17711 AF190051	Homo sapiens Gallus gallus	atopy related autoantigen CALC hepatocyte nuclear factor la	443	81
			dimerization cofactor isoform	277	
226 227	AK026256 Z69368	Homo sapiens Schizosacchar	unnamed protein product nuf2-like coiled-coil protein	866 230	98 25
		omyces pombe		11072	
228	AF275948	Homo sapiens	ABCA1	11763	99
229	AF161384	Homo sapiens	HSPC266	2006	98
230	Y16270	Homo sapiens	paralemin	1951	100 99
231	AJ245599	Homo sapiens	putative secreted ligand	2379 1545	99
232	W88499	Homo sapiens	Human stomach carcinoma clone HP10412-encoded protein.]
233	AF096286	Mus musculus	pecanex 1	3623	93
234	V64619_cd 1	Homo sapiens	30-NOV-1990 Human HE1 cDNA.	796	100
235	V64619_cd 1	Homo sapiens	30-NOV-1990 Human HE1 cDNA.	470	98
236	AF227258	Bos taurus	RPGR-interacting protein-1	1262	38
237	AJ132445	Homo sapiens	claudin-14	1181	100
238	AL034562	Homo sapiens	dJ684O24.2 (prodynorphin (Beta-	1330	100

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SEQ ID NO:	ACCESSION NUMBER	SPECIES	DESCRIPTION	SMITH- WATERMAN SCORE	MIDENTITY
			Neoendorphin-Dynorphin precursor, Proenkephalin B precursor))		
239	AF262027	Homo sapiens	eIF-5A2	808	100
240	AL079344	Arabidopsis · thaliana	putative protein	194	33
241	AC002394	Homo sapiens	Gene product with similarity to dynein beta subunit	1542	51
242	AJ271361	Takifugu rubripes	FRANK2 protein	303	30
243	AL021918	Homo sapiens	b34I8.1 (Kruppel related Zinc Finger protein 184)	1476	48
244	AF190167	Homo sapiens	membrane associated protein SLP-2	1736	99
245	Y10601	Homo sapiens	ankyrin-like protein	5877	100
246	AL121771	Homo sapiens	dJ548G19.1.1 (novel protein (ortholog of mouse zinc finger protein ZFP64) (translation of cDNA NT2RP3001398 (Em:AK001596)) (isoform 1))	3628	100
247	L25314	Drosophila melanogaster	actin-related protein	984	47
248	X63745	Homo sapiens	KDEL receptor	1095	100
249	AF112208	Homo sapiens	13kDa differentiation-associated protein	816	100
250	AP001707	Homo sapiens	human gene for claudin-8, Accession No. AJ250711	1172	100
251	AL136125	Homo sapiens	dJ304B14.1 (novel protein)	778	100
252	AL031186	Homo sapiens	bK984G1.1 (supported by FGENES)	532	100
253	Y17531	Homo sapiens	Human secreted protein clone BL205 14 protein.	639	100
254	AL049843	Homo sapiens	dJ392M17.3 (KIAA0349 protein)	6741	99
255	AJ242972	Homo sapiens	TOLLIP protein	1424	99
256	Y94873	Homo sapiens	Human protein clone HP02632.	1876	100
257	AF279865	Homo sapiens	kinesin-like protein GAKIN	2903	100
258 259	AL024498 R66278	Homo sapiens Homo sapiens	dJ417M14.1 (novel protein) Therapeutic polypeptide from glioblastoma cell line.	589 830	100
260	AF101784	Homo sapiens	b-TRCP variant E3RS-lkappaB	3226 .	99
261	AF101784	Homo sapiens	b-TRCP variant E3RS-IkappaB	2821	100
262	AF101784	Homo sapiens	b-TRCP variant E3RS-IkappaB	3149	99
263	AF197060	Homo sapiens	src homology 3 domain-containing protein HIP-55	2257	100
264	Y86262	Homo sapiens	Human secreted protein HAQAR23, SEQ ID NO:177.	766	100
265	Y56966.	Homo sapiens	Human SBPSAPL polypeptide.	2779	100
266	Y56966	Homo sapiens	Human SBPSAPL polypeptide.	1018	99
267	AJ300465	Homo sapiens	putative white family ATP-binding cassette transporter	1557	95
268	AC004030	Homo sapiens	F21856_2	3579	99
269	X55954	Homo sapiens	HL23 ribosomal protein	714	100
270	AB033921	Mus musculus	Ndr1 related protein Ndr2	1855	94
271	AF081886	Homo sapiens	ERO1-like protein	1905	99
272	AF166492	Homo sapiens	small GTPase RAB6B	1060	100
273 274	AL022238 W88667	Homo sapiens Homo sapiens	dJ1042K10.4 (novel protein) Secreted protein encoded by gene	2201 1530	99
05-	-		134 clone HAIBP89.		
275 276	X00129 Z47500_cd1	Homo sapiens Homo sapiens	precursor RBP 11-MAY-1998 Human RHOH gene	1044 1161	97 100
277	AB049188	Equus caballus	sequence. ubiquitin C-terminal hydrolase	1118	96

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No: SCORE	100 94 92 100 90 100 100 100
279 AF143956 Mus musculus coronin-2 2414 280 R85151 Homo sapiens Endothelial cell polypeptide. 911 281 R85151 Homo sapiens Endothelial cell polypeptide. 1031 282 D83948 Rattus S1-1 protein 3975 283 Y14768 Homo sapiens I Kappa B-like protein 2037 286 AL031316 Homo sapiens dJ28O10.3(HSD11B1 (hydroxysteroid (11-beta) dehydrogenase 1) 294 287 D64109 Homo sapiens MS4A7 1230 288 AB026043 Homo sapiens Krueppel-related DNA-binding protein 209 protein 290 AJ001810 Homo sapiens mRNA cleavage factor I 25 kDa subunit 1217 subunit 291 Y99454 Homo sapiens Human PRO1605 (UNQ786) amino acid sequence SEQ ID NO:395. 694	94 92 100 90 100 100
280 R85151 Homo sapiens Endothelial cell polypeptide. 911 281 R85151 Homo sapiens Endothelial cell polypeptide. 1031 282 D83948 Rattus S1-1 protein 3975 283 Y14768 Homo sapiens I Kappa B-like protein 2037 286 AL031316 Homo sapiens dJ28010.3(HSD11B1 (hydroxysteroid (11-beta) dehydrogenase 1) 294 287 D64109 Homo sapiens tob family 1773 288 AB026043 Homo sapiens Krueppel-related DNA-binding protein 209 290 AJ001810 Homo sapiens mRNA cleavage factor I 25 kDa subunit 1217 291 Y99454 Homo sapiens Human PRO1605 (UNQ786) amino acid sequence SEQ ID NO:395. 694	92 100 90 100 100
281 R85151 Homo sapiens Endothelial cell polypeptide. 1031 3975	100 90 100 100
282 D83948 Rattus S1-1 protein 3975	90 100 100 99 100
283 Y14768 Homo sapiens I Kappa B-like protein 2037	100 100 99 100
283 Y14768 Homo sapiens I Kappa B-like protein 2037 286 AL031316 Homo sapiens dJ28O10.3(HSD11B1 (hydroxysteroid (11-beta) dehydrogenase 1) 294 287 D64109 Homo sapiens tob family 1773 288 AB026043 Homo sapiens MS4A7 1230 289 M61866 Homo sapiens Krueppel-related DNA-binding protein 209 290 AJ001810 Homo sapiens mRNA cleavage factor I 25 kDa subunit 1217 291 Y99454 Homo sapiens Human PRO1605 (UNQ786) amino acid sequence SEQ ID NO:395. 694	99 100
286 AL031316 Homo sapiens dJ28O10.3(HSD11B1 294 (hydroxysteroid (11-beta) dehydrogenase 1) 1773 287 D64109 Homo sapiens tob family 1773 1230 288 AB026043 Homo sapiens MS4A7 1230 289 M61866 Homo sapiens Krueppel-related DNA-binding 209 protein 290 AJ001810 Homo sapiens mRNA cleavage factor I 25 kDa subunit 291 Y99454 Homo sapiens Human PRO1605 (UNQ786) amino acid sequence SEQ ID NO:395.	99 100
(hydroxysteroid (11-beta) dehydrogenase 1) 287	99 100
287 D64109 Homo sapiens tob family 1773 288 AB026043 Homo sapiens MS4A7 1230 289 M61866 Homo sapiens Krueppel-related DNA-binding protein 209 290 AJ001810 Homo sapiens mRNA cleavage factor I 25 kDa subunit 1217 291 Y99454 Homo sapiens Human PRO1605 (UNQ786) amino acid sequence SEQ ID NO:395. 694	100
288 AB026043 Homo sapiens MS4A7 1230 289 M61866 Homo sapiens Krueppel-related DNA-binding protein 209 290 AJ001810 Homo sapiens mRNA cleavage factor I 25 kDa subunit 1217 291 Y99454 Homo sapiens Human PRO1605 (UNQ786) amino acid sequence SEQ ID NO:395. 694	100
289 M61866 Homo sapiens Krueppel-related DNA-binding protein 290 AJ001810 Homo sapiens mRNA cleavage factor I 25 kDa subunit 291 Y99454 Homo sapiens Human PRO1605 (UNQ786) amino acid sequence SEQ ID NO:395.	
290 AJ001810 Homo sapiens mRNA cleavage factor I 25 kDa 1217 subunit 291 Y99454 Homo sapiens Human PRO1605 (UNQ786) amino acid sequence SEQ ID NO:395.	90
290 AJ001810 Homo sapiens mRNA cleavage factor I 25 kDa subunit 291 Y99454 Homo sapiens Human PRO1605 (UNQ786) amino acid sequence SEQ ID NO:395.	
291 Y99454 Homo sapiens Human PRO1605 (UNQ786) amino acid sequence SEQ ID NO:395.	100
acid sequence SEQ ID NO:395.	100
292 Y44824 Homo sapiens Human molecule associated with cell 2370	100
proliferation, MACP-4. 293 A1276101 Homo sapiens GPRC5B protein 2099	100
Zio illa illa illa illa illa illa illa il	100
27. 12.000 12.000 12.000 12.000	100
PRGE-21.	
296 U91561 Rattus pyridoxine 5'-phosphate oxidase 1239 norvegicus	87
297 L02956 Xenopus ribonucleoprotein 1624	83
298 AF226730 Homo sapiens Cyt19 1729	99
299 AF226730 Homo sapiens Cyt19 906	98
300 Y54324 Homo sapiens Amino acid sequence of a human 718	89
gastric cancer antigen protein.	
301 AF125533 Homo sapiens NADH-cytochrome b5 reductase 1606 isoform	100
302 Y32206 Homo sapiens Human receptor molecule (REC) 1676 encoded by Incyte clone 2825826.	98
303 AF247565 Homo sapiens hepatocellular carcinoma associated ring finger protein 525	100
304 AF208844 Homo sapiens BM-002 428	100
305 AC004983 Homo sapiens similar to PID:g3877944 1988	100
306 AL132978 Arabidopsis putative protein 210	25
307 Y10530 Homo sapiens olfactory receptor 1645	100
308 AF180681 Homo sapiens guanine nucleotide exchange factor 3597	100
309 AF111856 Homo sapiens sodium dependent phosphate 3591	99
transporter isoform NaPi-3b 310 Y13583 Homo sapiens G-protein coupled receptor 2171	100
310 Y 13383 Homo sapiens G-protein coupled receptor 2171 311 Z73420 Homo sapiens cE146D10.2 (mercaptopyruvate 1598	100
sulfurtransferase (EC 2.8.1.2))	
312 X79535 Homo sapiens beta tubulin 2348	100
313 AF070658 Homo sapiens HSPC002 861	100
314 AF078866 Homo sapiens SURF-4 1395	100
317 Z37986 Homo sapiens phenylalkylamine binding protein 1258	100
320 AB047892 Macaca hypothetical protein 258 fascicularis	82
321 Y25755 Homo sapiens Human secreted protein encoded 1440	100
from gene 45.	
	100 49

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SEQ	ACCESSION	SPECIES	DESCRIPTION	SMITH-	7 %
ID NO:	NUMBER		DESCRIPTION	WATERMAN SCORE	IDENTITY
		thaliana			
325	AF140501	Homo sapiens	DNA polymerase iota	3691	99
326	X96698	Homo sapiens	D1075-like	1450	96
327	AF152325	Homo sapiens	protocadherin gamma A5	4769	100
328	AF151803	Homo sapiens	CGI-45 protein	1970	100
329	X74070	Homo sapiens	transcription factor BTF3	639	81
330	AF171102	Homo sapiens	retinal degeneration B beta	1302	95
331	W54040	Homo sapiens	Human interferon-inducible protein, HIFI.	484	98
332	AF024617	Homo sapiens	transcription-associated zinc ribbon protein	691	100
333	U19181	Rattus norvegicus	Rabin3	2129	90
334	G03877	Homo sapiens	Human secreted protein, SEQ ID NO: 7958.	621	100
335	AL008582	Homo sapiens	bK223H9.2 (ortholog of A. thaliana F23F1.8)	626	100
336	AF110774	Homo sapiens	adrenal gland protein AD-001	647	100
337	AB011414	Homo sapiens	Kruppel-type zinc finger protein	1674	58
338	AF207600	Homo sapiens	ethanolamine kinase	129	100
340	AC020579	Arabidopsis thaliana	putative phosphoribosylformylglycinamidine synthase; 25509-29950	3283	50
341	Y28576	Homo sapiens	Secreted peptide clone pe503_1.	944	100
342	U32274	Saccharomyce s cerevisiae	Ydr386wp; CAI: 0.12	191	37
343	A01771	synthetic construct	vascular anticoagulating protein	1661	99
344	AF220052	Homo sapiens	uncharacterized hematopoietic stem/progenitor cells protein MDS032	1285	100
345	Y70400	Homo sapiens	Human cell-signalling protein-2.	754	100
346	Y50926	Homo sapiens	Human fetal brain cDNA clone vc16_1 derived protein.	962	100
347	AF183428	Homo sapiens	28.4 kDa protein	1329	100
348	AC006069	Arabidopsis	putative cleavage and	1383	55
		thaliana	polyadenylation specifity factor		
349	AL032631	Caenorhabditis elegans	Y106G6H.8	194	39
350	U70669	Homo sapiens	Fas-ligand associated factor 3	167	23
351	Y93468	Homo sapiens	Amino acid sequence of a potassium channel interactor protein.	1182	92
352	AF005856	Drosophila yakuba	anon2A5	111	45
353	AJ271684	Homo sapiens	myeloid DAP12-associating lectin	1013	100
354	AF099100	Homo sapiens	WD-repeat protein 6	2882	99
355	U51730	Murine leukemia virus	reverse transcriptase	316	42
356	D50617	Saccharomyce s cerevisiae	YFL042C	279	27
357	D50617	Saccharomyce s cerevisiae	YFL042C	279	27
358	AF161432	Homo sapiens	HSPC314	1059	93
359	AB029488	Homo sapiens	C11orf21	758	99
360	AJ251024	Homo sapiens	putative odorant binding protein ag	1239	100
361	U43281	Saccharomyce s cerevisiae	Lpg22p	2074	74
362	U43281	Saccharomyce s cerevisiae	Lpg22p	2153	74

SMITH-ACCESSION SPECIES DESCRIPTION SEQ NUMBER WATERMAN **IDENTITY** ID SCORE NO: Arabidopsis 100632 AC007153 156 24 363 thaliana AF197927 Homo sapiens AF5q31 protein 3992 99 364 Homo sapiens mitochondrial isoleucine tRNA 4286 98 365 D28500 synthetase 366 X97868 Homo sapiens arylsulphatase 3141 98 367 AL162048 Homo sapiens hypothetical protein 1532 100 L36062 Mus musculus steroidogenic acute regulatory 189 25 368 protein AF113249 Homo sapiens multiple domain putative nuclear 1022 59 369 protein endozepine-related protein precursor 2425 370 M15888 Bos taurus 84 X66363 Homo sapiens serine/threonine protein kinase 2562 100 371 372 W74802 Homo sapiens Human secreted protein encoded by 1532 89 gene 73 clone HSQEL25. tenascin-M1 AF100772 Homo sapiens 99 373 11535 PRO0518 374 AF090934 Homo sapiens 382 100 gonadotropin inducible transcription 375 AB021643 Homo sapiens 2761 99. repressor-3 1331 AB049758 MAWD binding protein 100 376 Homo sapiens Kruppel-associated box protein 377 AF070666 Homo sapiens 466 97 nuclear pore complex glycoprotein 464 60 S59342 Mus sp. 378 p62 Mus musculus Su(var)3-9 homolog Suv39h2 1690 88 379 AF149205 UDP-glucose:glycoprotein 380 AF227906 Homo sapiens 7851 99 glucosyltransferase 2 precursor hematopoietic zinc finger protein Mus musculus 1769 381 AF118566 92 unnamed protein product 382 AK000619 Homo sapiens 810 100 383 AF227906 Homo sapiens UDP-glucose:glycoprotein 7851 99 glucosyltransferase 2 precursor 384 AF117946 Link guanine nucleotide exchange 2363 100 Homo sapiens factor II 385 AF125390 Drosophila L82G 139 41 melanogaster 386 Y94907 Homo sapiens Human secreted protein clone 1092 50 ca106 19x protein sequence SEQ ID NO:20. 387 U18795 Saccharomyce Yel064cp 206 28 s cerevisiae 388 AF177388 Homo sapiens cancer-amplified transcriptional 10748 99 coactivator ASC-2 389 AJ002744 Homo sapiens UDP-GalNAc:polypeptide N-3469 96 acetylgalactosaminyltransferase 7 cone sodium-calcium potassium 390 AF097366 100 Homo sapiens 3166 exchanger 391 AF217525 Homo sapiens Down syndrome cell adhesion 5337 60 molecule 392 U81035 Rattus ankyrin binding cell adhesion 3967 91 norvegicus molecule neurofascin 393 X65224 Gallus gallus neurofascin 4097 78 X13916 Homo sapiens LDL-receptor related precursor (AA 4292 394 99 -19 to 4525) AF151083 Homo sapiens HSPC249 395 444 98 AB017026 Mus musculus oxysterol-binding protein 2173 396 98 397 AL035587 Homo sapiens dJ475N16.4 (KIAA0240) 2393 100 Human secreted protein encoded by 398 W74813 Homo sapiens 722 92 gene 85 clone HSDFV29. Human Hydrolase protein-8 399 Y71110 Homo sapiens 1637 99

(HYDRL-8).

	1 0 21 0 0 0 21 0 1 0 2

SEQ ID NO:	ACCESSION NUMBER	SPECIES	DESCRIPTION	SMITH- WATERMAN SCORE	MENTITY
400	AF039718	Caenorhabditis elegans	contains similarity to lupus LA protein homologs	325	43
401	AE000877	Methanotherm obacter thermoautotro phicus	conserved protein	231	36
402	Y27795	Homo sapiens	Human secreted protein encoded by gene No. 79.	1539	99
403	Z50853	Homo sapiens	CLPP	615	100
405	X03475	Rattus norvegicus	ribosomal protein L35a (aa 1-110)	:. 576	99
406	AF144237	Homo sapiens	LOMP protein	252	44
407	U20239	Mus musculus	fibrosin	288	76
409	AL033378	Homo sapiens	dJ323M4.1 (KIAA0790 protein)	6026	99
410	X54326	Homo sapiens	glutaminyl-tRNA synthetase	7577	99
411	X61585	Bos taurus	polynucleotide adenylyltransferase	3715	97
412	AF217190	Homo sapiens	MLEL1 protein	5271	99
414	G02815	Homo sapiens	Human secreted protein, SEQ ID NO: 6896.	314	95
415	AJ245922	Homo sapiens	alpha-tubulin 8	2370	100
416	AF203032	Homo sapiens	neurofilament protein	220	21
417	Z97653	Homo sapiens	c380A1.2.1 (novel protein (isoform 1))	1567	100
418	AJ404326	Homo sapiens	SR+89	1871	99
419	AJ404326	Homo sapiens	SR+89	902	64
420	AF134726	Homo sapiens	G9A	5334	99
421	L28125	Podospora anserina	beta transducin-like protein	288	39
422	W21733	Homo sapiens	NIP-1 encoded by clone 59.	110	72
423	S67970	Homo sapiens	ZNF75=KRAB zinc finger	951	76
424	L28035	Mus musculus	protein kinase C gamma	3768	98
426	Y73373	Homo sapiens	HTRM clone 921803 protein sequence.	555	56
427	Y73373	Homo sapiens	HTRM clone 921803 protein sequence.	266	49
428	X61118	Homo sapiens	TTG-2a/RBTN-2a	876	100
429	Z96932	Homo sapiens	nuclear autoantigen fo 14 kDa	496	83
430	AJ277291	Homo sapiens	HELG protein	678	72
431	X82157	Homo sapiens	hevin	3525	99
432	AC007192	Homo sapiens	P85B_HUMAN; PTDINS-3- KINASE P85-BETA	3825	99
433	AL021918	Homo sapiens	b34I8.1 (Kruppel related Zinc Finger protein 184)	1713	50
434	AF084464	Rattus norvegicus	GTP-binding protein REM2	141	29
435	AL049795	Homo sapiens	dJ622L5.2 (novel protein)	1756	98
436	M14513	Rattus norvegicus	(Na+ and K+) ATPase, alpha(III) catalytic subunit	4269	99
437	U33460	Homo sapiens	DNA-directed RNA polymerase I, largest subunit	8777	98
438	D87076	Homo sapiens	similar to human bromodomain protein BR140(JC2069)	3067	100
439	L43912	Macaca mulatta	mannose-binding protein A	589	93
440	D31763	Homo sapiens	ha0946 protein is Kruppel-related.	927	49
441	U70976	Homo sapiens	arrestin	2068	99
442	B08069	Homo sapiens	A human beta-alanine-pyruvate aminotransferase (HAPA).	2343	99
443	AF100662	Caenorhabditis	contains similarity to ubiquitin	166	24

SEQ ID NO:	ACCESSION NUMBER	SPECIES	DESCRIPTION	SMITH- WATERMAN SCORE	% IDENTITY
		elegans	carboxyl-terminal hydrolase (Pfam: UCH-1.hmm, score: 28.46) (Pfam: UCH-2.hmm, score: 47.53)		
444	D78017	Rattus norvegicus	NFI-A1	2667	98
445	AL049569	Homo sapiens	dJ37C10.3 (novel ATPase)	2418	100
448	AJ242540	Volvox carteri f. nagariensis	hydroxyproline-rich glycoprotein DZ-HRGP	165	34
449	AJ133352	Homo sapiens	ZNF237 protein	2006	100
450	AJ133352	Homo sapiens	ZNF237 protein	1025	96
451	AF170708	Homo sapiens	T-box protein TBX3	3700	99
452	AK002080	Homo sapiens	unnamed protein product	1546	99
453	L32977	Homo sapiens	Rieske Fe-S protein	1239	93
454	X51760	Homo sapiens	zinc finger protein (583 AA)	1533	57
455	Y01141	Homo sapiens	Secreted protein encoded by gene 7 clone HTLFA90.	1453	99
456	AB006631	Homo sapiens	The human homolog of mouse Cux-2	6559	100
457	AF067165	Homo sapiens	zinc finger protein 3	977	64
458	AF038169	Homo sapiens	unknown	154	38
459	W75214	Homo sapiens	Human secreted protein encoded by gene 19 clone HRSMC69.	1180	95
460	U97002	Caenorhabditis elegans	similar to acyl-CoA dehydrogenases and epoxide hydrolases; Pfam domain PF00441 (Acyl-CoA_dh), Score=57.4, E-value=1.7e-16, N=2; contains similarity to Pfam domain PF00702 (Hydrolase), Score=57.4, E-value=1e-13, N=1	583	37
461	AK023114	Homo sapiens	unnamed protein product	1041	99
462	M93134	Friend murine leukemia virus	pol protein	289	44
463	AF055473	Homo sapiens	GAGE-8	232	47
466	Y51415	Homo sapiens	Human wild type pKe83 protein.	2625	100
467	Y51417	787	Human pKe83 splice variant protein	2433	100
468	Y57936	Homo sapiens	Human transmembrane protein HTMPN-60.	1629	96
469	D38552	Homo sapiens	The ha1539 protein is related to cyclophilin.	2995	100
470	Y70013	Homo sapiens	Human Protease and associated protein-7 (PPRG-7).	3530	100
471	AJ224747	Homo sapiens	C-terminal variant of hINADL including 2 amino acid exchanges and an insertion of 28 amino acids in frame.	7969	100
472	W99665	Homo sapiens	Human secreted protein clone du 157_12 protein.	1546	100
473	W99665	Homo sapiens	Human secreted protein clone du157_12 protein.	998	98
474	X63526	Homo sapiens	homologue to elongation factor 1- gamma from A.salina	2273	99
475	X15940	Homo sapiens	ribosomal protein L31 (AA 1-125)	644	100
476	M60832	Homo sapiens	alpha-2 type VIII collagen	3581	99
477	AF039697	Homo sapiens	antigen NY-CO-31	1213	97
478	AF156929	Sus scrofa	inflammatory response protein 6	1588	83
479	AF264717	Homo sapiens	FYVE domain-containing dual specificity protein phosphatase FYVE-DSP2	5610	99
480	AF044578	Homo sapiens	putative DNA polymerase; POLAP	2478	94
481	X89750	Homo sapiens	TGIF protein	1413	100

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SEQ ID NO:	ACCESSION NUMBER	SPECIES	DESCRIPTION	SMITH- WATERMAN SCORE	% IDENTITY
482	M93107	Homo sapiens	(R)-3-hydroxybutyrate dehydrogenase	1663	96
483	U58334	Homo sapiens	Bbp/53BP2	1556	41
484	AF151538	Homo sapiens	deoxycytidyl transferase; Revlp	4281	99
485	Z98884	Homo sapiens	dJ467L1.1 (KJAA0833)	699	73
486	AJ243874	Homo sapiens	oligophrenin-4	3682	100
487	Z11737	Homo sapiens	flavin-containing monooxygenase 4	2969	100
488	X56123	Mus musculus	talin	4353	77
489	AJ278112	Homo sapiens	putative cell cycle control protein	335	23
490	W74843	Homo sapiens	Human secreted protein encoded by gene 115 clone HOVBA03.	1013	98
491	Y41337	Homo sapiens	Human secreted protein encoded by gene 30 clone HRDDV47.	509	36
492	X90530	Homo sapiens	ragB	1926	99
493	X90530	Homo sapiens	ragB	1405	99
494	X90530	Homo sapiens	ragB	1893	96
495	AL022394	Homo sapiens	dJ511B24.3 (KIAA0395 (probable homeobox protein))	4990	99
496	Y11395	Homo sapiens	lanthionine synthetase C-like protein 1	2168	100
497	AJ010119	Homo sapiens	Ribosomal protein kinase B (RSK-B)	4001	100
498	G01563	Homo sapiens	Human secreted protein, SEQ ID NO: 5644.	330	100
499	X54131	Homo sapiens	protein-tyrosine phosphatase	10465	99
500	G01082	Homo sapiens	Human secreted protein, SEQ ID NO: 5163.	549 3676	100
501	AC004142	Homo sapiens	similar to murine leucine-rich repeat protein; possible role in neural development by protein-protein interactions; 93% similarity to D49802 (PID:g1369906)	3070	100
502	AL117544	Homo sapiens	hypothetical protein	1226	100
503	AF203032	Homo sapiens	neurofilament protein	5115	99
504	AL034417	Homo sapiens	bK215D11.2 (similar to rat gene 33)	2476	100
505	X69090	Homo sapiens	190kD protein	7546	99
506	U58755	Caenorhabditis elegans	coded for by C. elegans cDNA yk34b1.5; coded for by C. elegans cDNA yk13h10.5; coded for by C. elegans cDNA yk46e8.5; coded for by C. elegans cDNA yk46d5.5; coded for by C. elegans cDNA yk43c2.5; coded for by C. elegans cDNA yk46e8.3; coded for by C. elegans cDNA yk43c2.3; coded for by C. elegans cDNA yk46d5.3; coded for by C. elegans cDNA yk13f10.3; coded for by C. elegans	782	55
507	AJ293309	Homo sapiens	cDNA yk34b1.3 NHP2 protein	801	100
508	U39045	Rattus norvegicus	cytoplasmic dynein intermediate chain 2B	3241	97
509	AF063231	Mus musculus	cytoplasmic dynein intermediate chain 2	3159	97
510	AF202893	Mus musculus	Kif21b	4336	95
511	Y13115	Homo sapiens	serine/threonine protein kinase	5071	99
512	AB030207	Homo sapiens	G gamma subunit	364	100
513	AF039571	Homo sapiens	peripheral benzodiazepine receptor interacting protein; PBR-IP/PRAX1	495	33
514	AB037883	Homo sapiens	Gb3/CD77 synthase	1916	99

SEQ ID NO:	ACCESSION NUMBER	SPECIES	DESCRIPTION	SMITH- WATERMAN SCORE	% IDENTITY
515	D90868	Escherichia coli	similar to	1489	100
516	X98834	Homo sapiens	zinc finger protein Hsal2	5290	100
517	AF055668	Mus musculus	apoptosis-linked gene 4, deltaC form	2904	78
518	AF019926	Mus musculus	protein kinase	1694	90
519	M34513	Homo sapiens	omega protein	317	91
520	Y08612	Homo sapiens	88kDa nuclear pore complex protein	2313	99
521	Y08612	Homo sapiens	88kDa nuclear pore complex protein	1561	99
522	AL096766	Homo sapiens	dA59H18.1 (KIAA0767 protein)	2497	100
523	AF186249	Homo sapiens	six transmembrane epithelial antigen of prostate	1790	100
524	AB029012	Homo sapiens	KIAA1089 protein	4933	100
525	AB026893	Homo sapiens	vascular cadherin-2	5962	100
526	X74331	Homo sapiens	DNA primase (p58 subunit)	1720	100
528	AC007228	Homo sapiens	R31665_2	1488	47
529	X14830	Homo sapiens	acetylcholine receptor beta-subunit preprotein	2639	100
530	U80446	Caenorhabditis elegans	coded for by C. elegans cDNA yk172e6.3; coded for by C. elegans cDNA yk158f7.3; coded for by C. elegans cDNA yk158f7.5; coded for by C. elegans cDNA yk172e6.5	420	39
531	S76838	Mus sp.	Dbs	4821	88
532	Z82215	Homo sapiens	dJ68O2.2 (myosin, heavy polypeptide 9, non-muscle)	9828	100
533	AF245505	Homo sapiens	adlican	277	31
534	AF300612	Homo sapiens	N-acetylgalactosamine-4-O- sulfotransferase	993	59
535	AL121928	Homo sapiens	bA18I14.3 (pleckstrin and Sec7 domain protein)	3333	99
536	AJ271055	Mus musculus	iroquois homeobox protein 6	1724	76
537	AF180473	Homo sapiens	Not2p	2267	100
538	AF071059	Mus musculus	zinc finger RNA binding protein	1089	. 51
539	AF023453	Homo sapiens	actin-related protein 3-beta	2219	100
540	AC003030	Homo sapiens	R29828_1	1401	70
541 542	AC003030 AL121889	Homo sapiens Homo sapiens	R29828_1 dJ1076E17.1 (KIAA0823 protein	2294 2152	100
		-	(continues in AL023803))		
543	AB006135	Rattus norvegicus	db83	1238	98
544	G02650	Homo sapiens	Human secreted protein, SEQ ID NO: 6731.	644	97
545	Y07595	Homo sapiens	transcription factor TFIIH	2373	100
546	AL133545	Homo sapiens	bA386N14.1 (novel protein similar to a dual specificity phosphatase)	964	99
547	X83618	Homo sapiens	hydroxymethylglutaryl-CoA synthase	2647	100
548	AF134726	Homo sapiens	NG37	4359	99
549	AB035356	Homo sapiens	neurexin I-alpha protein	6948	99
551	AB037901	Homo sapiens	gene amplified in squamous cell carcinoma-1	5215	99
552	AB043634	Homo sapiens	PAR-6A	885	100
553	AP000693	Homo sapiens	partial CDS	4875	99
554	AF002223	Homo sapiens	myotubularin related 1	3490	100
555	AC004893	Homo sapiens	similar to NEDD-4 (KIA0093); similar to P46934 (PID:g1171682)	1611	100
556	AJ404468	Homo sapiens	axonemal dynein heavy chain	8328	100
557	AJ404468	Homo sapiens	axonemal dynein heavy chain	11137	100

SEQ ID NO:	ACCESSION NUMBER	SPECIES	DESCRIPTION .	SMITH- WATERMAN SCORE	% IDENTITY
558	X65873	Homo sapiens	kinesin heavy chain	4860	100
559	AJ277365	Homo sapiens	polyglutamine-containing protein	592	36
560	AF205600	Homo sapiens	transposase-like protein	407	27
561	X71125	Homo sapiens	glutaminyl-peptide cyclotransferase	1914	100 .
562	X71125	Homo sapiens	glutaminyl-peptide cyclotransferase	1456	97
563	X54304	Homo sapiens	myosin regulatory light chain	897	100
564	AF250842	Drosophila melanogaster	multiple asters	130	23
565	Y58608	Homo sapiens	Protein regulating gene expression PRGE-1.	1619	99
566	AL121893	Homo sapiens	bA189K21.5 (novel protein similar to retinoblastoma binding protein (RBBP9))	1012	100
567	AL117352	Homo sapiens	dJ876B10.2 (novel protein (ortholog of rat EXO84))	3713	99
568	AF228603	Homo sapiens	pleckstrin 2	1841	100
569	AF239243	Homo sapiens	histone deacetylase 7	3244	86
570	AF087695	Mus musculus	veli 3	989	100
571	AB046381	Homo sapiens	testis-abundant finger protein	1346	99
572	AC005551	Homo sapiens	R26529_2, partial CDS	1020	100
573	Y90290	Homo sapiens	Human peptidase, HPEP-7 protein sequence.	274	52
574	W76734	Homo sapiens	Human mDia Rho targeting protein.	712	32
575	AL121935	Homo sapiens	bA517H2.3 (t-complex 10 (a murine tcp.homolog))	853	78
576	Y86217	Homo sapiens	Human secreted protein HWHGU54, SEQ ID NO:132.	2123	99
577	AL121716	Homo sapiens	dJ202D23.2 (novel protein)	6329	99
578	AL121716	Homo sapiens	dJ202D23.2 (novel protein)	6329	99
579	X92715	Homo sapiens	KRAB /C2H2 zinc finger protein	3102	97
580	X54637	Homo sapiens	protein tyrosine kinase	5564	98
581	X78817	Homo sapiens	p115	1148	44
582	AJ251245	Rattus norvegicus	SECIS binding protein 2	3086	71
583	AF113125	Homo sapiens	E-1 enzyme	581	100
584	M19529	Sus scrofa	follistatin A	1906	98
585	AF169677	Homo sapiens	leucine-rich repeat transmembrane protein FLRT3	3403	100
586	D87685	Homo sapiens	similar to human transcription factor TFIIS (S34159).	8083	99
587	Y00876	Homo sapiens	Human LAPH-1 protein sequence.	2110	100
588	Y99674	Homo sapiens	Human GTPase associated protein- 25.	2111	99
589	D86973	Homo sapiens	similar to Yeast translation activator GCN1 (P1:A48126)	12033	99
590	AL034452	Homo sapiens	dJ682J15.1 (novel Collagen triple helix repeat containing protein)	1979	100
591	¥57396	Homo sapiens	Human lysoenzyme LYC4 polypeptide.	814	100
592	AJ297743	Mus musculus	torsinB protein	1448	85
593	AF164796	Homo sapiens	NADH:ubiquinone oxidoreductase MLRQ subunit homolog	469	100
594	Y41312	Homo sapiens	Human secreted protein encoded by gene 5 clone HLDRM43.	749	94
595	Y41312	Homo sapiens	Human secreted protein encoded by	824	100
	1 11312		gene 5 clone HLDRM43.		<u> </u>
596	Y77123	Homo sapiens	gene 5 clone HLDRM43. Human neurotransmission-associated protein (NTAP) 998868.	2102	98

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SEQ ID NO:	ACCESSION NUMBER	SPECIES	DESCRIPTION	SMITH- WATERMAN SCORE	MIDENTITY
		melanogaster			
598	AF070447	Homo sapiens	barrier-to-autointegration factor	290	90
599	X56203	Plasmodium falciparum	liver stage antigen	372	22
600	X79828	Mus musculus	NK10	202	53
601	AB004109	Cricetulus griseus	phosphatidylserine synthase II	2262	92
602	U94988	Mus musculus	Nulp1	2912	89
603	U94988	Mus musculus	Nulp1	2800	86
604	AF006264	Homo sapiens	recombination and sister chromatid cohesion protein homolog	2850	100
605	AF006264	Homo sapiens	recombination and sister chromatid cohesion protein homolog	2530	100
606	X82260	Homo sapiens	RanGAP1	2929	100
607	X82260	Homo sapiens	RanGAP1	1843	97
608	AF160909	Drosophila melanogaster	BcDNA.LD03471	943	58
610	X74801	Homo sapiens	gamma subunit of CCT chaperonin	2745	99
611	AL031427	Homo sapiens	dJ167A19.1 (novel protein)	1608	100
612	Y71072	Homo sapiens	Human membrane transport protein, MTRP-17.	445	100
613	X16396	Homo sapiens	precursor polypeptide (AA -29 to 315)	1749	100
614	AK000281	Homo sapiens	unnamed protein product	1814	99
615	AB011128	Homo sapiens	KIAA0556 protein	5761	99
616	U19361	Petromyzon marinus	NF-180	205	21
617	AF045555	Homo sapiens	wbscr1	1208	100
618	AF045555	Homo sapiens	wbscr1 alternative spliced product	1318	100
619	U22229	Felis catus	ribosomal protein L41	128	100
620	Y17169	Homo sapiens	A6 related protein	1819	100
621	Y12065	Homo sapiens	hNop56	2956	99
622	AF177758	Homo sapiens	ubiquitin specific protease 16	2998	100
623	AF317425	Homo sapiens	GAC-1	3866	100
624	AL050297	Homo sapiens	hypothetical protein	1227	99
625	AC007204	Homo sapiens	BC273239_1	3398	99
626	Z68747	Homo sapiens	imogen 38	2024 1958	99 97
627 628	Z68747 Y70229	Homo sapiens	imogen 38 Human RNA-associated protein-10	3424	99
		Homo sapiens	(RNAAP-10).		
629	AF191492	Homo sapiens	nasopharyngeal carcinoma associated gene protein-8	613	100
630	AF119664	Homo sapiens	transcriptional regulator protein HCNGP	1574	100
631	AF119664	Homo sapiens	transcriptional regulator protein HCNGP	1150	89
632	Y17849	Homo sapiens	ganglioside-induced differentiation associated protein 1	1839	98
633	X55740	Homo sapiens	5'-nucleotidase	3012	100
634	AF03968°	Homo sapiens	antigen NY-CO-3	931	100
635	AF119662	Homo sapiens	E46 protein	2424	100
636	AB007836	Homo sapiens	Hic-5	2544	100
637	AF077818	Mus musculus	syntrophin-associated serine- threonine protein kinase	2027	44
638	AL035455	Homo sapiens	dJ1018E9.1 (VAMP (vesicle- associated membrane protein)- associated protein B and C)	150	26
639	AF078844	Homo sapiens	hqp0376 protein	416	81

SEQ ID NO:	ACCESSION NUMBER	SPECIES	DESCRIPTION	SMITH- WATERMAN SCORE	mentity
640	U28377	Escherichia coli	ORF_f239; was ORF_f191 and ORF_f194 before splice	1198	100
641	AK024442	Homo sapiens	FLJ00032 protein	1677	56
642	U58682	Homo sapiens	ribosomal protein S28	340	100
643	X57432	Rattus rattus	ribosomal protein S2	1520	98
644	AB002348	Homo sapiens	KIAA0350 protein	5186	. 99
646	Y96202	Homo sapiens	IkappaB kinase (IKK) binding protein, Y2H56.	1178	98 ,
647	AB029482	Mus musculus	JNK-binding protein JNKBP1	4609	81
648	AB009053	Arabidopsis thaliana	contains similarity to isoamyl acetate-hydrolyzing esterase—gene_id:MQB2.25	407	44
650	AC002550	Homo sapiens	Unknown gene product	858	99
651	U26592	Homo sapiens	diabetes mellitus type I autoantigen	253	66
652	X60155	Homo sapiens	zinc finger 41	4349	100
653	X53330	Platynereis dumerilii	H4 protein (AA 1 - 103)	523	100
654	AC003682	Homo sapiens	R27945_2	2558	100
655	X80473	Mus musculus	rab19	596	56
656	J02649	Rattus norvegicus	unknown protein	201	95
657	AC006014	Homo sapiens	similar to RFP transforming protein; similar to P14373 (PID:g132517)	1331	99
658	X92972	Homo sapiens	protein phosphatase 6	1666	100
659	L35269	Homo sapiens	zinc finger protein	2803	99
660	AC003682	Homo sapiens	F18547_1	3184	96
661	X79204	Homo sapiens	ataxin-1	4195	99
662	X17620	Homo sapiens	Nm23 protein	965	99
663	AB015617	Homo sapiens	ELKS	1501	80
664	Z56281	Homo sapiens	interferon regulatory factor 3	2331	100
665	AJ248283	Pyrococcus abyssi	LACTOYLGLUTATHIONE LYASE (EC 4.4.1.5) METHYLGLYOXALASE) (ALDOKETOMUTASE) (GLYOXALASE I).	254	40
666	Z70200	Homo sapiens	U5 snRNP-specific 200kD protein	8819	99
667	Z70200	Homo sapiens	U5 snRNP-specific 200kD protein	8589.	97
668	AF153450	Manduca sexta	juvenile hormone esterase binding protein	225	32
669	AF227198	Homo sapiens	CrkRS	7231	99
670	X99586	Homo sapiens	SMT3C protein	441	87
671	Z61589_cd1	Homo sapiens	17-AUG-1998 DNA encoding a human OC-2 protein.	2593	100
672	AJ132702	Mus musculus	ATFa-associated factor	3240	88
673	AF204159	Homo sapiens	potassium large conductance calcium-activated channel beta 3a subunit	1486	100
674	G02061	Homo sapiens	Human secreted protein, SEQ ID NO: 6142.	558	99
675	G01246	Homo sapiens	Human secreted protein, SEQ ID NO: 5327.	141	77
676	AB016839	Homo sapiens	mob1	419	42
677	D86970	Homo sapiens	similar to myosin heavy chain: Containing ATP/GTP-binding site motif A(P-loop)	161	28
678	U83115	Homo sapiens	non-lens beta gamma-crystallin like protein	. 8569	99
679	AF203687	Homo sapiens	prolactin regulatory element-binding protein	2181	100

SEQ ID NO:	ACCESSION NUMBER	SPECIES	DESCRIPTION	SMITH- WATERMAN SCORE	% IDENTITY
680	M27685	Mus musculus	ultra-high sulphur keratin	650	58
681	U04968	Cricetulus griseus	nucleotide excision repair protein	3712	97
682	AF119663	Homo sapiens	G-protein gamma-12 subunit	356	100
683	G03733	Homo sapiens	Human secreted protein, SEQ ID NO: 7814.	342	100
684	X67699	Homo sapiens	CDw52 antigen	297	100
685	AF022789	Homo sapiens	ubiquitin hydrolyzing enzyme I	1892	100
686	AJ001006	Mus musculus	EMeg32 protein	938	96
687	W03516	Homo sapiens	Prostaglandin DP receptor.	1864	100
688	AF019661	Mus musculus	zeta proteasome chain; PSMA5	1214	100
689	AF156557	Homo sapiens	stomatin related protein	2036	100
690	G03960	Homo sapiens	Human secreted protein, SEQ ID NO: 8041.	593	100
691	AF161512	Homo sapiens	HSPC163	738	100
692	AL031115	Homo sapiens	ZXDA, ZXDB (zinc finger X-linked protein)	4298	100
693	L40410	Homo sapiens	thyroid receptor interactor	806	100
694	AC004542	Homo sapiens	OXYSTEROL-BINDING PROTEIN-like; similar to P22059 (PID:g129308)	2533	99
695	AF169411	Rattus norvegicus	PAPIN	4144	52
696	Y58168	Homo sapiens	Human hydrolase homologue HHH-4.	2144	100
697	AF271994	Homo sapiens	dopamine responsive protein DRG-1	1613	100
698	Y41741	Homo sapiens	Human PRO704 protein sequence.	1323	100
699	AL133506	Unknown	/prediction=(method:""genscan"", version:""1.0"", score:""109.13""); /prediction=(method:	825	48
700	Y96870	Homo sapiens	Human goose-type lysozyme (GOLY).	1032	100
701	AC003034	Homo sapiens	Gene with similarity to rat kidney- specific (KS) gene	1190	100
702	AC003034	Homo sapiens	Gene with similarity to rat kidney- specific (KS) gene	937	95
703	AJ242832	Homo sapiens	calpain	3756	100
704	S52624	Homo sapiens	unknown	185	100
705.	AF005081	Homo sapiens	skin-specific protein	652	100
706	·Y16793	Homo sapiens	keratin, type I	2232	100
707	Y44985	Homo sapiens	Human epidermal protein-2.	455	69
708	AF113220	Homo sapiens	MSTP040	686	100
709	Y44985	Homo sapiens	Human epidermal protein-2.	408	65
710	Y16132	Homo sapiens	CDT6	1874	100 .
711	Y68775	Homo sapiens	Amino acid sequence of a human phosphorylation effector PHSP-7.	2407	100
712	X63422	Homo sapiens	H(+)-transporting ATP synthase	209	100
713	AF169968	Mus musculus	DNA binding protein DESRT	1467	79
714	X52563	Bos taurus	permability increasing protein	383	29
715	AJ277739	Homo sapiens	RPB11b1alpha protein	480	98
716	AL135791	Homo sapiens	bA162G10.3 (zinc finger protein)	401	98
717	AF223466	Homo sapiens	HT015 protein	1311	97
719	AF117383	Homo sapiens	placental protein 13; PP13	746	100
720	Z98743	Homo sapiens	dJ181C9.2 (Rho GTPase activating protein 8 (RhoGAP, p50RhoGAP))	324	100
721	AL163815	Arabidopsis thaliana	putative protein	653	61
722	G01436	Homo sapiens	Human secreted protein, SEQ ID	418	96

SEQ	ACCESSION	SPECIES	DESCRIPTION	SMITH-	%
ID NO:	NUMBER	ļ		WATERMAN SCORE	IDENTITY
110.			NO: 5517.	SCORE	
723	AF282919	Mus musculus	Zfp228	349	49
724	AB023191	Homo sapiens	KIAA0974 protein	2953	100
725	AL031778	Homo sapiens	dJ34B21.1 (novel BZRP	920	100
			(benzodiazapine receptor (peripheral)		
			(MBR, PBR, PBKS, IBP, Isoquinoline-binding protein)) LIKE		[
			protein)		
726	AL021939	Homo sapiens	dJ352A20.2 (aldehyde	1764	100
	1 2023,00	IIIIII Supremo	dehydrogenase family protein)	.,	100
727	AF182426	Rattus	arylacetamide deacetylase	791	42
		norvegicus			
728	Y08565	Homo sapiens	UDP-GalNAc:polypeptide N-	3331	99
	15155105		acetylgalactosaminyltransferase	1.000	
729	AF155135	Homo sapiens	novel retinal pigment epithelial cell protein	1652	99
730	AL078606	Arabidopsis	putative protein	277	55
1 730	/LU/6000	thaliana	putative protein	211)))
731	Y73352	Homo sapiens	HTRM clone 1732368 protein	1720	100
			sequence.		
732	AF178432	Homo sapiens	SH3 protein	3302	100
733	Y17832	Human	env protein	223	34
		endogenous retrovirus K			
734	Y28859	Homo sapiens	Human mesoderm induction early	2067	98
'54	120037	Tionio supiciis.	response protein ER1.	2007	90
735	U09355	Oryctolagus	protein phosphatase 2A1 B gamma	2352	99
		cuniculus	subunit		
736	Y94922	Homo sapiens	Human secreted protein clone pv6_1	724	99
707	A TO 0 500 0		protein sequence SEQ ID NO:50.		
737	AB027003 AF112200	Mus musculus Homo sapiens	protein phosphatase NADH-oxidoreductase B18 subunit	378 739	84
739	AF112200	Homo sapiens	NADH-oxidoreductase B18 subunit	613	100 88
740	AF302154	Homo sapiens	SPG protein	6556	100
741	B25681	Homo sapiens	Human secreted protein sequence	1410	99
		•	encoded by gene 17 SEQ ID NO:70.		
742	L27479	Homo sapiens	X123	1237	99
743	L27479	Homo sapiens	X123	1206	97
744	Y66745	Homo sapiens	Membrane-bound protein PRO1186.	588	99
745 746	AJ001019 X68453	Homo sapiens - Sus scrofa	ring finger protein	1292	99
747	Y57897	Homo sapiens	tubulin-tyrosine ligase Human transmembrane protein	1882 1173	94 100
'''	13,03,	liomo suprens	HTMPN-21.	1175	100
748	AF151069	Homo sapiens	HSPC235	1694	96
749	AF182404	Homo sapiens	mitochondrial uncoupling protein 1	1674	100
750	AL121993	Homo sapiens	dJ776P7.1 (Novel protein)	2500	99
751	AF149825	Homo sapiens	PACSIN3	2253	100
752	AL008635	Homo sapiens	dJ510H16.2 (high-mobility group	3026	99
753	Y57914	Homo sapiens	protein 2-like 1) Human transmembrane protein	1124	100
661	13/714	riomo sabiens	HTMPN-38.	1124	100
754	AF285109	Homo sapiens	septin 3 isoform B	1766	100
755	AF004161	Oryctolagus	peroxisomal Ca-dependent solute	2371	95
		cuniculus	carrier		
756	Z19585	Homo sapiens	thrombospondin-4	4239	100
757	AP001745	Homo sapiens	similar to zinc finger 5 protein	1857	100
758 759	AF190664 AF090326	Mus musculus Mus musculus	LMBR2	555	72
760	AF090326 AL096677	Homo sapiens	AE-1 binding protein AEBP2 dJ322G13.3 (novel protein similar to	1540 999	97 94
	111070011	Tromo advicus	work of the state	777	74

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SEQ ID NO:	ACCESSION NUMBER	SPECIES	DESCRIPTION	SMITH- WATERMAN SCORE	MENTITY
			bovine and mouse beta-soluble NSF attachment protein (SNAP-beta))		
761	AC003007	Homo sapiens	Unknown gene product (partial)	649	96
762	U66372	Bos taurus	ribosomal protein S29	230	73
764	Y90899	Homo sapiens	D1-like dopamine receptor activity modifying protein SEQ ID NO:1.	1152	100
765	U88169 .	Caenorhabditis elegans	similar to molybdoterin biosynthesis MOEB proteins	1204	65
766	AL118506	Homo sapiens	dJ591C20.3.1 (novel DnaJ domain protein, similar to mouse and bovine cysteine string protein)	1091	100
767	AK024693	Homo sapiens	unnamed protein product	3767	100
768	Z11518	Homo sapiens	histidyl-tRNA synthetase	2582	100
769	X13916	Homo sapiens	LDL-receptor related precursor (AA -19 to 4525)	25529	100
770	AC009360	Arabidopsis thaliana	Contains 3 PF 00400 WD40, G-beta repeat domains.	333	33
771	AB037685	Mus musculus	LANP-like protein	1246	91
772	AL161578	Arabidopsis thaliana	putative protein	335	46
773	AL161578	Arabidopsis thaliana	putative protein	333	47
774	AY008271	Homo sapiens	helicase SMARCAD1	5264	99
775	Y21591	Homo sapiens	Human secreted protein (clone CC332-33).	1127	96
776	W88853	Homo sapiens	Polypeptide fragment encoded by gene 89.	752	100
777	W88853	Homo sapiens	Polypeptide fragment encoded by gene 89.	752	100
778	W88853	Homo sapiens	Polypeptide fragment encoded by gene 89.	752	100
779	AF196481	Homo sapiens	RING finger protein; FXY2	3644	100
780	AL035427	Homo sapiens	dJ769N13.1 (KIAA0443 protein.)	1609	54
781	AB026187	Homo sapiens	protocadherin-Xa	5244	100
782	B24458	Homo sapiens	Human secreted protein sequence encoded by gene 22 SEQ ID NO:83.	1002	100
783	AB027289	Homo sapiens	cyclin-E binding protein 1	5421	100
784	G02916	Homo sapiens	Human secreted protein, SEQ ID NO: 6997.	627	100
785	AJ245822	Homo sapiens	type I transmembrane receptor	4560	100
786	AJ245820	Homo sapiens	type I transmembrane receptor	4624 3340	100 99
787 788	Z48042 AL031782	Homo sapiens Homo sapiens	GPI-anchored protein p137 dJ708F5.1 (PUTATIVE novel	2739	100
			Collagen alpha 1 LIKE protein)		100
789	AJ131245	Homo sapiens	Sec24B protein	6602	100
790	AF107203	Homo sapiens	ataxin 2-binding protein	2008 600	100
791 792	Y14690 AL031055	Homo sapiens Homo sapiens	procollagen alpha 2(V) dJ28H20.2 (novel protein)	1267	34 100
792	Y36194	787	Human secreted protein	2051	99
794	AB028127	Homo sapiens	mannosyltransferase	2138	96
795	AC007228	Homo sapiens	R31665 2	2738	79
796	AL049482	Arabidopsis thaliana	putative protein	436	47
797	AC004528	Homo sapiens	R32184 3	891	91
798	AB037830	Homo sapiens	KIAA1409 protein	7532	100
799	X53793	Homo sapiens	5' half of the product is homologues to Bacillus subtiis SAICAR synthetase, 3' half corresponds to the catalytic subunit of AIR carboxylase	2232	100

	YAIYIACY				
SEQ ID NO:	ACCESSION NUMBER	SPECIES	DESCRIPTION	SMITH- WATERMAN SCORE	% IDENTITY
800	Y99350	Homo sapiens	Human PRO1378 (UNQ715) amino acid sequence SEQ ID NO:33.	1343	100
801	AB042636	Homo sapiens	junctophilin type3	1225	47
802	AB029324	Rattus norvegicus	TIP120-family protein TIP120B	3916	90
803	AB029324	Rattus norvegicus	TIP120-family protein TIP120B	4961	90
804	AF251040	Homo sapiens	putative nuclear protein	2119	100
805	AB033281	Homo sapiens	F-box and WD-repeats protein beta- TRCP2 isoform C	2879	100
806	U87305	Rattus norvegicus	transmembrane receptor UNC5H1	3257	90
807	AF118889	Rattus norvegicus	b-tomosyn isoform	3155	97
808	AF226993	Rattus norvegicus	selective LIM binding factor	8793	95
809	W19919	Homo sapiens	Human Ksr-1 (kinase suppressor of Ras).	3939	99
810	AL031782	Homo sapiens	dJ708F5.1 (PUTATIVE novel Collagen alpha 1 LIKE protein)	1546	100
811	AC002542	Homo sapiens	similar to C. elegans F11A10.5; 80% similarity to Z68297 (PID:g1130619)	2294	100
812	U83246	Homo sapiens	copine I	606	52
813	AF242552	Gallus gallus	retinovin .	945	34
814	X52332	Homo sapiens	zinc finger protein 10	1651	93
815	X52332	Homo sapiens	zinc finger protein 10	2423	99
816	Y09631	Homo sapiens	PIBF1 protein	2935	99
817	X71997	Rattus norvegicus	myosin I	3883	98
818	AY004877	Mus musculus	cytoplasmic dynein heavy chain	11105	98
819	Y27196	Homo sapiens	Human cyclic nucleotide phosphodiester PDE8B(E) amino acid sequence.	3790	100
820	AF081947	Mus musculus	tektin	1134	81
821	AL035106	Homo sapiens	dJ998C11.1 (continues in Em:AL445192 as bA269H4.1)	871	100
822	AF022795	Homo sapiens	TGF beta receptor associated protein-	385	24
823	AF015770	Mus musculus	radical fringe	1422	82
824	U82695	Homo sapiens	expressed-Xq28STS protein	1444	99
825	X77371	Mesocricetus auratus	COR1	641	78
826	AB014576	Homo sapiens	KIAA0676 protein	296	79
827	AL049733	Homo sapiens	dJ875H3.1 (APK1 antigen)	1584	72
828	AF222980	Homo sapiens	disrupted in Schizophrenia 1 protein	4418	100
829	Z31560	Homo sapiens	sox-2	1683	100
830	AF295773	Homo sapiens	ral guanine nucleotide dissociation stimulator	4717	99
831	AB041926	Homo sapiens	GCK family kinase MINK-2	6866	100
832	L04948	Saccharomyce s cerevisiae	mitochondrial transporter protein	338	35
833	AJ007012	Mus musculus	Fish protein	704	94
834	Z34289	Homo sapiens	nucleolar phosphoprotein p130	3455	99
835	U10991	Homo sapiens	G2	8436	98
836	AF230877	Homo sapiens	MIP-T3	2945	99
837	X58288	Homo sapiens	protein-tyrosine phosphatase	7734	99
838	X56958	Homo sapiens	ankyrin (brank-2)	9631	100
839	AC024791	Caenorhabditis elegans	contains similarity to beta-lactamases	370	24

SEQ	ACCESSION	SPECIES	DESCRIPTION	· SMITH-	%
ID NO:	NUMBER		1	WATERMAN SCORE	IDENTITY
840	D83197	Homo sapiens	ankyrin repeat protein	802	99
841	AF053711	Serinus canaria	neurofilament medium subunit	192	31
842	AF283772	Homo sapiens	similar to Homo sapiens ribosomal protein L10 encoded by GenBank Accession Number L25899	990	96
843	U76343	Homo sapiens	GABA transport protein	2992	98
844	Y13645	Homo sapiens	uroplakin II	897	100
845	D21064	Homo sapiens	similar to rat general mitochondrial matrix processing protease mRNA (RATMPP).	2710	99
846	AF192522	Homo sapiens	Niemann-Pick C3 protein; NPC3	7047	100
847	AF192522	Homo sapiens	Niemann-Pick C3 protein; NPC3	5472	100
848	X60489	Homo sapiens	elongation factor-1-beta	1162	100
849	AC007204	Homo sapiens	BC273239_1	2277	67
850	AC003682	Homo sapiens	R28830_1	2401 353	100
851 852	AL121583 Z48475	Homo sapiens Homo sapiens	bA358N2.1 (novel protein) glucokinase regulator	3155	61 99
853	Z83844	Homo sapiens	dJ37E16.2 (SH3-domain binding protein 1)	1884	98
854	AF233323	Homo sapiens	Fas-associated phosphatase-1	390	36
855	AF062741	Rattus norvegicus	pyruvate dehydrogenase phosphatase isoenzyme 2	447	80
856	Y11411	Homo sapiens	pristanoyl-CoA oxidase	3595	98
857	M97188	Strongylocentr otus purpuratus	tektin A1	290	46
858	AB001105	Homo sapiens	hippocalcin-like protein 4	995	100
859	AF164791	Homo sapiens	putative 38.3kDa protein	1795	100
860	AF298117	Homo sapiens	homeobox protein OTX2	1477	93
861	AF015264	Rattus norvegicus	golgi peripheral membrane protein p65	1820	81
862	X16901	Homo sapiens	30kb subunit of RAB30 /74	1284	100 .
863	M12140	Homo sapiens	envelope protein	202	81
864	AF161459	Homo sapiens	HSPC109	815	98
865	AL109983	Homo sapiens	dJ718P11.1.1 (novel class II aminotransferase similar to serine palmotyltransferase (isoform 1))	444	100
866	M77183	Rattus norvegicus	alpha-1-macroglobulin	227	45
867	AF272663	Homo sapiens	gephyrin	3785	100
868	X75285	Mus musculus	fibulin-2	3258	87
869	X82494	Homo sapiens	fibulin-2	3407	99
870	AJ297743	Mus musculus	torsinB protein	169	43
871	AJ278313	Homo sapiens	phospholipase C-beta-1a	6258	99
872 873	AF073344 Y91955	Homo sapiens Homo sapiens	ubiquitin-specific protease 3 Human cytoskeleton associated protein 10 (CYSKP-10).	256 535	100
874	AJ000414	Homo sapiens	Cdc42-interacting protein 4	1136	53
875	AF265555	Homo sapiens	ubiquitin-conjugating BIR-domain enzyme APOLLON	627	100
876	Y48586	Homo sapiens	Human breast tumour-associated protein 47.	2537	98
877	AF182198	Homo sapiens	intersectin 2 long isoform	8764	99
878	L17308	Gossypium hirsutum	proline-rich cell wall protein	192	35
879	AF177169	Homo sapiens	tropomodulin 2	1769	100
880	W03627	Homo sapiens	Human follicle stimulating hormone	210	23
			GPR N-terminal sequence.		

SEQ	ACCESSION	SPECIES	DESCRIPTION	SMITH-	%
ID NO:	NUMBER			WATERMAN SCORE	IDENTITY
881	AL021068	Homo sapiens	dJ206D15.3	2615	99
882	AC005498	Homo sapiens	R31665_2	318	82
883	AF165518	Homo sapiens	MAGOH isoform	182	94
884	D21211	Homo sapiens	protein tyrosine phosphatase (PTP-BAS, type 3)	368	43
885	U13045	Homo sapiens	nuclear respiratory factor-2 subunit beta 1	869	62
886	X52836	Homo sapiens	tryptophan hydroxylase (AA 1 - 444)	2320	98
887	X51466	Homo sapiens	elongation factor 2	4460	100
888	AB039903	Homo sapiens	interferon-responsive finger protein 1 long form	1096	98
889	X51760	Homo sapiens	zinc finger protein (583 AA)	3130	100
890	AJ243396	Homo sapiens	voltage-gated sodium channel beta-3 subunit	1024	100
891	W67928	Homo sapiens	Fragment of human secreted protein encoded by gene 4.	391	100
892	AB020598	Homo sapiens	peptide transporter 3	3017	100
893	Y66648	Homo sapiens	Membrane-bound protein PRO1120.	4722	99
894	Y66648	Homo sapiens	Membrane-bound protein PRO1120.	3606	96
895	A29218 cd	Homo sapiens	19-NOV-1998 DNA encoding G-	2178	100
	1		protein coupled 7 TM receptor with AXOR15 activity.		
896	AJ000332	Homo sapiens	Glucosidase II	5063	99
897	X98259	Homo sapiens	M-phase phosphoprotein 8	1085	100
898	X57110	Homo sapiens	c-cbl protein	4849	99
899	X63652	Homo sapiens	inter-alpha-trypsin inhibitor heavy chain ITIH1	3376	98
900	X85134	Homo sapiens	RB protein binding protein	2816	99
901	L11672	Homo sapiens	zinc finger protein	2047	58
902	Y85565	Homo sapiens	Human homologue of UNC-53 (Hs-UNC-53/2) sequence.	369	83
903	X54871	Homo sapiens	ras related protein Rab5b	1094	100
904	Z98265	Homo sapiens	plakophilin 3	4065	100
905	AL035295	Homo sapiens	hypothetical protein	959	99
906	AF051782	Homo sapiens	diaphanous 1	801	35
907	AF208536	Homo sapiens	nucleotide binding protein; NBP	1372	100
908	U79240	Homo sapiens	serine/threonine protein kinase	2365.	. 98
909	U79240	Homo sapiens	serine/threonine protein kinase	2386	99
910	AJ132545	Homo sapiens	protein kinase	2921	100
911	AJ132545	Homo sapiens	protein kinase	1637	99
912	AL121733	Homo sapiens	hypothetical protein	1344	99
913	Y67579	Homo sapiens	Human death inducer-obliterator 1 (DIO-1) polypeptide.	1586	100
914	X87342	Homo sapiens	Human giant larvae homologue	5317	99
915	X87342	Homo sapiens	Human giant larvae homologue	3495	96
916	M94362	Homo sapiens	lamin B2	2357	93
917	AJ011654	Homo sapiens	triple LIM domain protein	3432	100
918	AJ131899	Rattus norvegicus	proline rich synapse associated protein 1	5776	88
919	AF054986	Homo sapiens	putative transmembrane GTPase	1816	100
920	U95822	Homo sapiens	putative transmembrane GTPase	1237	100
921	Y11588	Homo sapiens	apoptosis specific protein	1492	100
922	X84195	Homo sapiens	acylphosphatase	510	100
923	U72882	Homo sapiens	interferon-induced leucine zipper protein	1409	99
924	AE000660	Homo sapiens	ĥADV36S1	573	100
925	AF126245	Homo sapiens	acyl-Coenzyme A dehydrogenase-8	2162	100
	<u> </u>	<u></u>	precursor		

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SEQ ID NO:	ACCESSION NUMBER	SPECIES	DESCRIPTION	SMITH- WATERMAN SCORE	MENTITY
926	AE001968	Deinococcus radiodurans	hypothetical protein	147	27
927	W81576	Homo sapiens	EBV-induced G-protein coupled receptor (EBI-2) polypeptide.	1778	100
928	U01317	Homo sapiens	beta-globin	687	94
929	X98333	Homo sapiens	organic cation transporter	2933	100
930	Y91444	Homo sapiens	Human secreted protein sequence encoded by gene 42 SEQ ID NO:165.	1401	100
931	Y91644	Homo sapiens	Human secreted protein sequence encoded by gene 43 SEQ ID NO:317.	1243	100
932	D90279	Homo sapiens	collagen alpha 1(V) chain precursor	569	39
933	Z31560	Homo sapiens	sox-2	1587	96
934	AF147790	· Homo sapiens	transmembrane mucin 12	3047	99
935	Z85996	Homo sapiens	match: multiple proteins; match: Q08151 P28185 Q01111 Q43554; match: Q08150 Q40195 P20340 Q39222; match: Q40368 P36412 P40393 Q40723; match: CE01798 Q38923 Q40191 Q41022; match: Q39433 Q40177 Q40218 Q08146; match: P10949 P11023 Q16948 Q20337; match: Q25389 P25228 P20336 P05713; match: P35276 Q08147 P17609 P22128; match: Q15771 P36410 P35291; GTP- binding	726	94
936	AB041533	Homo sapiens	sperm antigen	1054	38
937	X91906	Homo sapiens	voltage-gated chloride ion channel	3914	100
938	AB032481	Homo sapiens	homeobox transcription factor	1744	100
939	AF111106	Homo sapiens	protein serine/threonine phosphatase 4 regulatory subunit 1	4682	99
940	Y17999	Homo sapiens	Dyrk1B protein kinase	3331	99
941	AF305872	Homo sapiens	thyroglobulin	455	92
942	AF263462	Homo sapiens	cingulin	5939	99
943	AK024442	Homo sapiens	FLJ00032 protein	1616	61
944	Y35911	Homo sapiens	Extended human secreted protein sequence, SEQ ID NO. 160.	262	35
945	AB015320	Homo sapiens	sigma1B subunit of AP-1 clathrin adaptor complex	599	71
946	Z82287	Caenorhabditis elegans	ZK550.2	229	35
947	D84223	Homo sapiens	leucyl tRNA synthetase	6207	99
948	U49057	Rattus norvegicus	rA9	3846	62
949	AK000568	Homo sapiens	unnamed protein product	1659	100
950	AL021578	Homo sapiens	dJ453C12.6.1 (uncharacterized hypothalamus protein (isoform 1))	257	42
951	AB032435	Homo sapiens	differentiation-associated Na- dependent inorganic phosphate cotransporter	3063	99
952	AF110532	Homo sapiens	uncoupling protein UCP-4	1561	100
953	X83587	Mus musculus	1A13 protein	1420	59
954	AL031665	Homo sapiens	dJ545L17.5.1 (novel protein)	386	53
955	Y87600	Homo sapiens	Human fatty acid synthase-like protein (HFASLP).	2377	100
956	Y99421	Homo sapiens	Human PRO1433 (UNQ738) amino acid sequence SEQ ID NO:292.	522	55

SEQ ID NO:	ACCESSION NUMBER	SPECIES	DESCRIPTION	SMITH- WATERMAN SCORE	% IDENTITY
957	U68535	Mus musculus	aldo-keto reductase	451	73
958	AC007067	Arabidopsis thaliana	T10O24.10	1594	57
959	U72194	Mus musculus	muskelin	3947	99
960	AE003661	Drosophila melanogaster	CG15168 gene product	277	54
961	X80332	Mus musculus	rab20	983	82
962	Y67315	Homo sapiens	Human secreted protein BL89_13 amino acid sequence.	3916	99
963	Y67315	Homo sapiens	Human secreted protein BL89_13 amino acid sequence.	3916	99
964	L32602	Rattus norvegicus	homeodomain 159341	1821	96
965	Z97832	Homo sapiens	dJ329A5.3 (KIAA06460 protein)	3581	99
966	W88995	Homo sapiens	Polypeptide fragment encoded by gene 146.	176	39
967	U12465	Homo sapiens	ribosomal protein L35	604	100
968	AF151803	Homo sapiens	CGI-45 protein	1101	78
969	W74865	Homo sapiens	Human secreted protein encoded by gene 137 clone HMWIF35.	1348	98
970	L21936	Homo sapiens	succinate dehydrogenase flavoprotein subunit	703	100
971	AJ133521	Drosophila buzzatii	protease, reverse transcriptase, ribonuclease H, integrase	194	23
972	AC006017	Homo sapiens	N-acetylgalactosaminyltransferase; similar to Q10473 (PID:g1709559)	3271	100
973	Z81317	Schizosacchar omyces pombe	DNA2-NAM7 helicase family protein	685	31
974	M17885	Homo sapiens	acidic ribosomal phosphoprotein (P0)	792	100
975	U22829	Mus musculus	P2Y purinoceptor	399	40
976	AL132772	Homo sapiens	dJ1013A22.1 (hepatic nuclear factor 4, alpha)	2466	99
977	AC003973	Homo sapiens	ZNF91L	1550	43
978	J04031	Homo sapiens	MDMCSF (EC 1.5.1.5; EC 3.5.4.9; EC 6.3.4.3)	2824	63
979	AF136715	Homo sapiens	taxol resistant associated protein	217	76
980	AF136715	Homo sapiens	taxol resistant associated protein	306	95
981	Z92822	Caenorhabditis elegans	ZK520.1	1109	44
982	AJ295149	Homo sapiens	putative dipeptidase	1564	99
983	AL021331	Homo sapiens	dJ366N23.3 (KIAA0173 and Tubulin-Tyrosine Ligase LIKE)	1492	100
984	AL161501	Arabidopsis thaliana	putative adenosine deaminase	370	38

TABLE 3

SEQ ID NO:	ACCESSION NO.	DESCRIPTION .	RESULTS*
2	BL00282	Kazal serine protease inhibitors family proteins.	BL00282 16.88 4.259e-14 97-120
3	BL00298	Heat shock hsp90 proteins family proteins.	BL00298A 10.97 1.000e-40 74- 119 BL00298E 27.30 1.000e-40 321-376 BL00298F 11.21 1.000e- 40 409-464 BL00298H 20.50 1.000e-40 553-607 BL00298C 16.40 2.286e-40 186-230

SEQ ID NO:	ACCESSION NO.	DESCRIPTION	RESULTS*
			BL00298B 15.64 1.290e-39 134- 181 BL00298G 24.57 5.345e-39 465-520 BL00298I 30.07 7.818e- 34 661-715 BL00298D 17.97 6.226e-33 242-282
4	PR00237	RHODOPSIN-LIKE GPCR SUPERFAMILY SIGNATURE	PR00237A 11.48 4.316e-13 57-82
5	PD02454	!!!! PROTEIN ALU SUBFAMILY WARNING ENTRY NUCLEAR PHOSPHO.	PD02454B 11.61 4.309e-17 75- 103
6	DM00864	EGF-LIKE DOMAIN.	DM00864A 15.21 7.429e-09 98- 119
7	PR00237	RHODOPSIN-LIKE GPCR SUPERFAMILY SIGNATURE	PR00237A 11.48 1.750e-11 29-54 PR00237D 8.94 7.000e-09 138- 160 PR00237B 13.50 8.250e-09 61-83
9	PF00855	PWWP domain proteins.	PF00855 13.75 5.667e-15 272-289
10	BL00139	Eukaryotic thiol (cysteine) proteases cysteine proteins.	BL00139D 9.24 4.400e-11 391- 408 BL00139A 10.29 7.511e-09 67-77
12	BL01113	C1q domain proteins.	BL01113B 18.26 9.294e-19 689- 725 BL01113C 13.18 4.857e-11 757-777 BL01113D 7.47 2.161e- 10 790-800
13	BL01113	C1q domain proteins.	BL01113B 18.26 3.813e-14 599-635 BL01113C 13.18 4.857e-11 667-687 BL01113D 7.47 2.161e-10 700-710
14	BL00594	Aromatic amino acids permeases proteins.	BL00594A 16.75 6.531e-10 50-94
15	BL01047	Heavy-metal-associated domain proteins.	BL01047B 19.73 4.913e-13 707- 728
16	PR00625	DNAJ PROTEIN FAMILY SIGNATURE	PR00625A 12.84 7.462e-18 310- 330 PR00625B 13.48 3.939e-15 340-361
18	BL00615	C-type lectin domain proteins.	BL00615A 16.68 3.700e-09 144- 162
20	PR00741	GLYCOSYL HYDROLASE FAMILY 29 SIGNATURE	PR00741D 16.11 9.082e-21 175- 195 PR00741F 14.66 9.262e-21 243-265 PR00741B 14.23 1.947e- 18 128-145 PR00741G 9.29 2.180e-17 318-340 PR00741C 9.16 7.328e-17 147-166 PR00741H 10.32 2.141e-13 351- 374 PR00741A 9.24 3.596e-13 89-105 PR00741E 13.39 3.535e- 12 215-232
22	BL00107	Protein kinases ATP-binding region proteins.	BL00107A 18.39 3.647e-20 117- 148 BL00107B 13.31 1.000e-16 182-198
23	BL00107	Protein kinases ATP-binding region proteins.	BL00107A 18.39 1.600e-23 126- 157
24	BL00107	Protein kinases ATP-binding region proteins.	BL00107A 18.39 1.600e-23 126- 157
27	BL00239	Receptor tyrosine kinase class II proteins.	BL00239B 25.15 2.324e-16 91- 139
28	BL00018	EF-hand calcium-binding domain proteins.	BL00018 7.41 3.250e-10 681-694 BL00018 7.41 6.400e-10 717-730
29	BL00018	EF-hand calcium-binding domain	BL00018 7.41 3.250e-10 681-694

11 34 9			
SEQ	ACCESSION	DESCRIPTION	RESULTS*
D	NO.		
NO:		·	
		proteins.	BL00018 7.41 6.400e-10 717-730
30	.BL01113	Clq domain proteins.	BL01113A 17.99 9.308e-09 54-81
33	PD01168	SYNTHETASE LIGASE PROTEIN	PD01168L 9.47 1.667e-09 401-
		ALANYL.	416
34	PD01168	SYNTHETASE LIGASE PROTEIN ALANYL.	PD01168L 9.47 1.667e-09 411- 426
36	PR00426	C5A-ANAPHYLATOXIN RECEPTOR SIGNATURE	PR00426D 10.59 3.618e-12 110- 122
37	PF00791	Domain present in ZO-1 and Unc5-like netrin receptors.	PF00791B 28.49 2.049e-10 1080- 1135
38	BL00350	MADS-box domain proteins.	BL00350 20.79 1.000e-40 1-55
40	BL00330	Alkaline phosphatase proteins.	BL00123B 19.31 1.000e-40 90-
40	BL00123	Alkainie phosphatase proteins.	133 BL00123C 24.61 1.000e-40 145-195 BL00123E 22.25 1.000e- 40 304-358 BL00123G 26.01 1.000e-40 438-488 BL00123F 19.03 8.714e-35 364-399 BL00123A 10.80 9.000e-24 52-77 BL00123D 12.73 1.000e-17 216- 229
44	PD00066	PROTEIN ZINC-FINGER METAL- BINDI.	PD00066 13.92 2.800e-14 346-359 PD00066 13.92 4.600e-14 486-499 PD00066 13.92 1.000e-13 374-387 PD00066 13.92 6.000e-13 458-471 PD00066 13.92 2.714e-12 234-247 PD00066 13.92 3.143e-12 430-443 PD00066 13.92 8.714e-12 514-527 PD00066 13.92 3.739e-11 402-415 PD00066 13.92 2.038e-10 318-331
45	DM00973	3 kw RESISTANCE BENOMYL YLL028W CYCLOHEXIMIDE.	DM00973A 21.17 2.946e-10 180- 217
47	BL00649	G-protein coupled receptors family 2	BL00649C 17.82 1.682e-10 475-
47	BL00049	proteins.	501 BL00649B 20.68 7.387e-09 417-463
50	PD00066	PROTEIN ZINC-FINGER METAL-BINDI.	PD00066 13.92 8.200e-16 445-458 PD00066 13.92 5.846e-15 305-318 PD00066 13.92 1.000e-14 221-234 PD00066 13.92 1.000e-14 417-430 PD00066 13.92 2.800e-14 249-262 PD00066 13.92 2.800e-14 277-290 PD00066 13.92 8.800e-14 333-346 PD00066 13.92 9.400e-14 361-374 PD00066 13.92 4.000e-13 389-402 PD00066 13.92 6.571e-12 473-486
51	BL00226	Intermediate filaments proteins.	BL00226D 19.10 1.000e-40 417- 464 BL00226B 23.86 3.348e-35 251-299 BL00226C 13.23 1.429e- 24 316-347 BL00226A 12.77 1.857e-15 151-166
52	PR00217	43 KD POSTSYNAPTIC PROTEIN SIGNATURE	PR00217C 10.91 5.648e-09 133- 149
53	BL00232	Cadherins extracellular repeat proteins domain proteins.	BL00232B 32.79 1.000e-40 143- 191 BL00232A 27.72 2.350e-28 49-82 BL00232B 32.79 7.052e-21 252-300 BL00232C 10.65 6.625e- 20 250-268 BL00232B 32.79 1.314e-11 367-415 BL00232C 10.65 9.308e-10 470-488

S-100/ICaBP type calcium binding

54

BL00303

BL00303B 26.15 8.759e-23 125-

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SEQ ID NO:	ACCESSION NO.	DESCRIPTION	RESULTS*
		protein.	162 BL00303A 21.77 1.000e-21 82-119
58	PR00378	INOSITOL PHOSPHATASE SIGNATURE	PR00378D 16.86 1.000e-15 242- 261 PR00378B 13.80 9.250e-13 109-129
59	PR00425	BRADYKININ RECEPTOR SIGNATURE	PR00425C 13.23 9.040e-12 120- 140
60	BL00280	Pancreatic trypsin inhibitor (Kunitz) family proteins.	BL00280 24.61 6.727e-38 238-282 BL00280 24.61 1.514e-30 294-338
65	BL01019	ADP-ribosylation factors family proteins.	BL01019A 13.20 1.222e-11 43-83
68	PR00237	RHODOPSIN-LIKE GPCR SUPERFAMILY SIGNATURE	PR00237E 13.03 5.091e-13 188- 212 PR00237G 19.63 7.207e-13 268-295 PR00237A 11.48 4.375e- 11 24-49 PR00237C 15.69 3.057e-10 101-124 PR00237D 8.94 4.750e-10 137-159 PR00237F 13.57 5.364e-10 230- 255 PR00237B 13.50 9.438e-10 57-79
70	PD01066	PROTEIN ZINC FINGER ZINC- FINGER METAL-BINDING NU.	PD01066 19.43 7.938e-28 31-70
71	PR00830	ENDOPEPTIDASE LA (LON) SERINE PROTEASE (S16) SIGNATURE	PR00830A 8.41 8.759e-12 348- 368
72	BL00120	Lipases, serine proteins.	BL00120B 11.37 2.149e-10 148- 163
77	PR00753	1-AMINOCYCLOPROPANE-1- CARBOXYLATE SYNTHASE SIGNATURE	PR00753E 8.01 3.552e-11 191- 216 PR00753D 6.85 2.778e-09 131-153
78	PR00506	D21 CLASS N6 ADENINE-SPECIFIC DNA METHYLTRANSFERASE SIGNATURE	PR00506C 19.40 8.017e-09 96- 119
82	BL00107	Protein kinases ATP-binding region proteins.	BL00107A 18.39 3.571e-16 436- 467
84	BL00675	Sigma-54 interaction domain proteins ATP-binding region A proteins.	BL00675A 24.86 8.800e-10 256- 300
85	BL00027	'Homeobox' domain proteins.	BL00027 26.43 2.286e-30 117-160
87	BL00250	TGF-beta family proteins.	BL00250A 21.24 6.786e-36 264- 300 BL00250B 27.37 1.450e-26 328-364
91	BL00215	Mitochondrial energy transfer proteins.	BL00215A 15.82 9.250e-17 10-35 BL00215A 15.82 6.000e-16 221- 246 BL00215A 15.82 7.857e-12 108-133 BL00215B 10.44 9.526e- 11 168-181
92	BL00027	'Homeobox' domain proteins.	BL00027 26.43 9.526e-24 324-367
95	PR00094	ADENYLATE KINASE SIGNATURE	PR00094C 12.94 1.000e-08 119-
96	PD02327	GLYCOPROTEIN ANTIGEN PRECURSOR IMMUNOGLO.	PD02327B 19.84 2.091e-09 143- 165
97	BL00752	XPA protein.	BL00752B 19.17 7.309e-09 28-72
98	PR00876	NEMATODE METALLOTHIONEIN SIGNATURE	PR00876B 7.66 2.268e-10 135- 149
99	PR00109	TYROSINE KINASE CATALYTIC DOMAIN SIGNATURE	PR00109B 12.27 9.824e-12 122- 141
100	BL00027	'Homeobox' domain proteins.	BL00027 26.43 7.429e-31 118-161
101	BL00028	Zinc finger, C2H2 type, domain proteins.	BL00028 16.07 6.870e-12 370-387 BL00028 16.07 6.885e-11 398-415 BL00028 16.07 8.269e-11 342-359 BL00028 16.07 4.300e-10 229-246

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SEQ	ACCESSION	DESCRIPTION	RESULTS*
ID	NO.		
NO:			DI 00028 16 07 6 1000 10 258 275
102	PR00048	C2H2-TYPE ZINC FINGER	BL00028 16.07 6.100e-10 258-275 PR00048A 10.52 7.750e-14 665-
102	1 100046	SIGNATURE	679 PR00048A 10.52 8.500e-14
	· ·		581-595 PR00048A 10.52 9.250e-
			14 637-651 PR00048A 10.52
			2.059e-12 609-623 PR00048A 10.52 2.588e-12 469-483
		}	PR00048A 10.52 7.353e-12 553-
1			567 PR00048A 10.52 2.895e-11
1			525-539 PR00048A 10.52 4.316e-
			11 441-455 PR00048A 10.52 5.263e-11 413-427 PR00048B
			6.02 2.125e-10 569-579
			PR00048B 6.02 4.938e-10 513-
			523 PR00048A 10.52 5.696e-10
			497-511 PR00048B 6.02 8.875e- 10 429-439 PR00048B 6.02
			1.000e-09 457-467 PR00048B
			6.02 6.684e-09 485-495
103	PR00195	DYNAMIN SIGNATURE	PR00195A 11.94 5.364e-22 31-50
			PR00195B 9.47 1.783e-21 56-74 PR00195C 11.50 3.455e-21 126-
			144 PR00195D 11.76 8.714e-21
1			175-194 PR00195F 16.20 8.500e-
		·	20 217-237 PR00195E 9.82
104	BL01113	C1q domain proteins.	8.650e-20 194-211 BL01113A 17.99 1.865e-09 121-
104	BEOTTIS	Crq domain proteins.	148 BL01113A 17.99 5.846e-09
			82-109
105	BL00420	Speract receptor repeat proteins domain proteins.	BL00420A 20.42 6.400e-11 70-99 BL00420A 20.42 8.525e-10 73-
		proteins.	102 BL00420A 20.42 5.708e-09
L			85-114
108	PR00860	VERTEBRATE METALLOTHIONEIN	PR00860B 7.04 2.929e-20 27-41
		SIGNATURE	PR00860A 5.46 5.500e-16 5-18 PR00860C 9.61 1.474e-14 41-51
112	BL01031	Heat shock hsp20 proteins family profile.	BL01031C 17.68 6.400e-10 122-
			147
114	DM01840	kw SPAC24B11.09 R07E5.13.	DM01840B 22.04 2.688e-40 59-
			103 DM01840A 10.95 9.571e-13 31-43
115	BL01126	Elongation factor Ts proteins.	BL01126A 18.48 2.317e-30 46-89
			BL01126B 13.15 7.387e-19 116-
			135 BL01126C 9.20 9.735e-11 190-203
116	BL00216	Sugar transport proteins.	BL00216B 27.64 4.375e-21 35-85
118	BL00437	Catalase proximal heme-ligand proteins.	BL00437A 18.82 1.000e-40 49-
			101 BL00437B 16.28 1.000e-40
			114-168 BL00437C 21.86 1.000e- 40 190-239 BL00437D 25.72
			1.000e-40 248-301 BL00437E
			23.95 1.000e-40 327-379
119	BL00140	Ubiquitin carboxyl-terminal hydrolase	BL00140D 22.64 8.274e-14 164-
		family 1 cysteine activ.	208 BL00140C 11.80 5.444e-10 77-102
120	BL00224	Clathrin light chain proteins.	BL00224B 16.94 6.712e-10 95-
	-2002		148
122	BL00203	Vertebrate metallothioneins proteins.	BL00203 13.94 1.000e-40 16-62
123	PR00041	CAMP RESPONSE ELEMENT	PR00041D 7.95 2.906e-09 24-41

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SEQ ID NO:	ACCESSION NO.	DESCRIPTION	RESULTS*
		BINDING (CREB) PROTEIN SIGNATURE	
124	PR00041	CAMP RESPONSE ELEMENT BINDING (CREB) PROTEIN SIGNATURE	PR00041D 7.95 2.906e-09 24-41
125	BL00061	Short-chain dehydrogenases/reductases family proteins.	BL00061C 7.86 3.250e-10 212- 222
126	PD01066	PROTEIN ZINC FINGER ZINC- FINGER METAL-BINDING NU.	PD01066 19.43 6.400e-25 251-290
127	PR00318	ALPHA G-PROTEIN (TRANSDUCIN) SIGNATURE	PR00318D 16.28 1.900e-34 219- 248 PR00318B 14.79 3.455e-27 168-191 PR00318C 12.09 7.000e- 23 197-215 PR00318A 7.84 1.600e-19 35-51 PR00318E 7.23 2.500e-12 265-275
128	PR00927	ADENINE NUCLEOTIDE TRANSLOCATOR 1 SIGNATURE	PR00927E 14.93 9.743e-10 67-89 PR00927B 14.66 4.575e-09 69-91
130	BL00824	Elongation factor 1 beta/beta/delta chain proteins.	BL00824B 9.21 7.750e-22 133- 153
131	BL00824	Elongation factor 1 beta/beta'/delta chain proteins.	BL00824C 14.58 1.000e-40 166- 204 BL00824D 14.04 1.621e-38 204-239 BL00824B 9.21 7.750e- 22 133-153 BL00824E 12.49 1.000e-19 247-263
132	PR00209	ALPHA/BETA GLIADIN FAMILY SIGNATURE	PR00209B 4.88 9.222e-13 1209- 1228
133	PR00209	ALPHA/BETA GLIADIN FAMILY SIGNATURE	PR00209B 4.88 9.222e-13 1168- 1187
134	PR00708	ALPHA-1-ACID GLYCOPROTEIN SIGNATURE	PR00708D 14.67 1.000e-27 141- 168 PR00708C 11.77 1.643e-25 98-120 PR00708B 15.15 2.174e- 24 73-95 PR00708E 13.33 1.600e-21 189-207 PR00708A 14.40 2.636e-21 51-70
135	PR00109	TYROSINE KINASE CATALYTIC DOMAIN SIGNATURE	PR00109B 12.27 8.468e-13 126- 145
136	PF00023	Ank repeat proteins.	PF00023A 16.03 3.250e-10 201- 217
137	BL00471	Small cytokines (intercrine/chemokine) C-x-C subfamily signat.	BL00471 23.92 7.480e-10 42-90
140	PR00205	CADHERIN SIGNATURE	PR00205B 11.39 5.582e-10 328- 346 PR00205B 11.39 9.018e-10 543-561
141	BL00412	Neuromodulin (GAP-43) proteins.	BL00412D 16.54 7.704e-09 976- 1027
143	PR00979	TAFAZZIN SIGNATURE	PR00979E 10.83 5.950e-26 192- 214 PR00979A 11.91 8.773e-25 63-83 PR00979C 12.16 6.400e-19 108-124 PR00979D 12.38 7.955e- 19 170-185 PR00979F 10.14 3.382e-15 230-244 PR00979B 15.59 5.636e-15 94-106
145	DM00686	kw REPLICATION REP 28K 17.7K.	DM00686C 14.14 7.720e-09 111- 131
146	PR00604	CLASS IA AND IB CYTOCHROME C SIGNATURE	PR00604D 15.86 1.000e-17 87- 104 PR00604B 12.73 9.591e-16 57-73 PR00604C 10.21 8.200e-12 73-84 PR00604E 10.13 1.000e-11 106-117 PR00604A 11.13 8.800e-

SEQ	ACCESSION	DESCRIPTION	RESULTS*
ID NO:	NO.	DESCRIPTION	
			11 44-52 PR00604F 8.60 1.000e- 10 123-132
147	BL00107	Protein kinases ATP-binding region proteins.	BL00107A 18.39 3.864e-15 266- 297 BL00107B 13.31 6.143e-11 335-351
148	PD00289	PROTEIN SH3 DOMAIN REPEAT PRESYNA.	PD00289 9.97 8.448e-09 67-81
149	PR00069	ALDO-KETO REDUCTASE SIGNATURE	PR00069D 19.36 1.857e-30 187- 217 PR00069A 16.01 7.429e-25 41-66 PR00069E 18.14 3.100e-22 235-260 PR00069C 16.03 7.000e- 20 151-169 PR00069B 11.33 8.071e-19 101-120
150	BL00027	'Homeobox' domain proteins.	BL00027 26.43 2.688e-27 139-182
151	PD02906	SYNTHASE I PSEUDOURIDYLATE PSEUDOURIDINE LYASE TR.	PD02906C 24.17 7.070e-22 165- 200 PD02906B 15.35 8.393e-15 114-127 PD02906A 10.84 6.500e- 09 71-84
153	BL00479	Phorbol esters / diacylglycerol binding domain proteins.	BL00479A 19.86 5.091e-12 891- 914 BL00479B 12.57 1.837e-11 915-931
158	BL00027	'Homeobox' domain proteins.	BL00027 26.43 6.786e-31 143-186
160	BL00422	Granins proteins.	BL00422C 16.18 7.750e-12 420- 448
162	PR00625	DNAJ PROTEIN FAMILY SIGNATURE	PR00625A 12.84 9.297e-11 62-82
164	BL01282	BIR repeat proteins.	BL01282B 30.49 6.182e-10 347- 386
166	PR00860	VERTEBRATE METALLOTHIONEIN SIGNATURE	PR00860B 7.04 2.929e-20 83-97 PR00860A 5.46 1.000e-18 61-74 PR00860C 9.61 1.900e-15 97-107
167	PR00449	TRANSFORMING PROTEIN P21 RAS SIGNATURE	PR00449A 13.20 7.052e-09 196- 218
169	BL00514	Fibrinogen beta and gamma chains C-terminal domain proteins.	BL00514C 17.41 1.346e-39 316- 353 BL00514G 15.98 2.241e-34 471-501 BL00514H 14.95 6.571e- 27 510-535 BL00514E 14.28 1.273e-16 388-405 BL00514D 15.35 9.100e-15 369-382 BL00514B 16.42 4.857e-14 260- 276 BL00514F 11.65 9.690e-14 416-431 BL00514A 11.68 8.200e- 11 149-159
170	BL00514	Fibrinogen beta and gamma chains C-terminal domain proteins.	BL00514C 17.41 1.346e-39 268- 305 BL00514G 15.98 2.241e-34 423-453 BL00514H 14.95 6.571e- 27 462-487 BL00514E 14.28 1.273e-16 340-357 BL00514D 15.35 9.100e-15 321-334 BL00514B 16.42 4.857e-14 212- 228 BL00514F 11.65 9.690e-14 368-383 BL00514A 11.68 8.200e- 11 101-111
171	BL00514	Fibrinogen beta and gamma chains C-terminal domain proteins.	BL00514G 15.98 2.241e-34 385- 415 BL00514H 14.95 6.571e-27 424-449 BL00514C 17.41 4.632e- 24 230-267 BL00514E 14.28 1.273e-16 302-319 BL00514D 15.35 9.100e-15 283-296

SEQ ID NO:	ACCESSION NO.	DESCRIPTION	RESULTS*
NO.			BL00514B 16.42 4.857e-14 212- 228 BL00514F 11.65 9.690e-14 330-345 BL00514A 11.68 8.200e-
1.77	DI 00007	ITTo-manhard domain metains	11 101-111 BL00027 26.43 9.400e-29 119-162
173 174	BL00027 DM01970	'Homeobox' domain proteins. 0 kw ZK632.12 YDR313C	DM01970B 8.60 5.119e-15 1391-
		ENDOSOMAL III.	1404
176	BL00773	Chitinases family 19 proteins.	BL00773C 9.42 8.000e-09 2-16
182	PR00109	TYROSINE KINASE CATALYTIC DOMAIN SIGNATURE	PR00109B 12.27 9.163e-14 141- 160
183	PD01937	DNA PROTEIN POLYMERASE ENDONUCLEASE DNA	PD01937A 6.68 3.475e-09 221- 232
185	BL00845	CAP-Gly domain proteins.	BL00845 16.43 2.946e-23 247-272 BL00845 16.43 1.628e-21 107-132
186	PR00452	SH3 DOMAIN SIGNATURE	PR00452B 11.65 6.538e-11 525- 541
187	PR00452	SH3 DOMAIN SIGNATURE	PR00452B 11.65 6.538e-11 497- 513
188	DM01803	1 HERPESVIRUS GLYCOPROTEIN H.	DM01803A 10.51 1.000e-09 1081-1102
189	PF00651	BTB (also known as BR-C/Ttk) domain proteins.	PF00651 15.00 5.091e-15 69-82
190	PR00194	TROPOMYOSIN SIGNATURE	PR00194C 6.38 1.900e-35 145- 174 PR00194E 8.74 3.250e-30 231-257 PR00194D 9.57 1.500e- 26 175-199 PR00194B 10.24 5.200e-24 120-141 PR00194A 7.86 4.857e-21 84-102
192	PD02042	IRON-SULFUR ELECTRON TRANSPORT AROMATIC HYDROCARB.	PD02042B 16.75 5.154e-09 131- 146 PD02042A 21.13 5.909e-09 94-121
193	PR00021	SMALL PROLINE-RICH PROTEIN SIGNATURE	PR00021A 4.31 2.200e-10 2-15
195	BL00463	Fungal Zn(2)-Cys(6) binuclear cluster domain proteins.	BL00463 8.22 5.071e-09 111-123
196	PR00118	BETA-LACTAMASE CLASS A SIGNATURE	PR00118F 16.42 9.386e-09 165- 181
197	DM00215	PROLINE-RICH PROTEIN 3.	DM00215 19.43 5.424e-09 234- 267
198	BL00660	Band 4.1 family domain proteins.	BL00660A 31.50 5.500e-11 714- 767
199	BL00282	Kazal serine protease inhibitors family proteins.	BL00282 16.88 8.820e-13 70-93
202	PR00009	TYPE I EGF SIGNATURE	PR00009A 14.15 5.345e-15 971- 987 PR00009C 14.11 8.773e-13 996-1008 PR00009D 16.83 8.000e-11 1008-1018 PR00009C 14.11 1.882e-09 892-904
203	BL00025	P-type 'Trefoil' domain proteins.	BL00025 17.17 4.536e-19 38-59
205	BL00018	EF-hand calcium-binding domain proteins.	BL00018 7.41 7.300e-10 165-178
206	PR00168	SLOW VOLTAGE-GATED POTASSIUM CHANNEL SIGNATURE	PR00168D 12.88 6.865e-11 67-86
207	BL00025	P-type 'Trefoil' domain proteins.	BL00025 17.17 3.423e-20 39-60 BL00025 17.17 8.750e-16 88-109
209	BL00646	Ribosomal protein S13 proteins.	BL00646B 21.42 6.100e-30 110- 143 BL00646A 25.82 6.192e-29 14-62
210	PR00138	MATRIXIN SIGNATURE	PR00138D 16.56 3.605e-25 279-

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SEQ	NO.	DESCRIPTION	MESOUIS.
NO:			
			305 PR00138C 16.41 3.000e-24
			218-247 PR00138E 6.01 8.714e-
			13 314-328 PR00138A 15.14
			9.538e-13 134-148 PR00138B 15.82 4.522e-12 188-204
211	DM01206	CORONA VIRUS NUCLEOCAPSID	DM01206B 10.69 8.429e-12 386-
211	DW101200	PROTEIN.	406 DM01206B 10.69 1.247e-10
		TROTEIN.	384-404 DM01206B 10.69
	·		5.068e-10 388-408
212	PD01941	TRANSMEMBRANE	PD01941A 14.81 1.000e-40 163-
	·	COTRANSPORTER SYMP.	217 PD01941B 15.02 9.705e-30
			420-467 PD01941E 15.92 8.714e-
			23 837-884 PD01941C 19.96
ļ		1	8.200e-20 508-563 PD01941D
			27.18 1.600e-16 661-710 PD01941F 28.52 9.645e-15 1005-
			1060
213	BL00362	Ribosomal protein S15 proteins.	BL00362 24.67 8.313e-09 330-373
214	BL00302 BL00115	Eukaryotic RNA polymerase II	BL00115Z 3,12 2.125e-09 1178-
		heptapeptide repeat proteins.	1227 BL00115Z 3.12 6.096e-09
			1164-1213
215	BL00038	Myc-type, 'helix-loop-helix' dimerization	BL00038B 16.97 7.600e-18 125-
		domain proteins.	146 BL00038A 13.61 1.474e-13
			102-118
216	BL01108	Ribosomal protein L24 proteins.	BL01108A 20.33 2.241e-22 49-82
	*		BL01108B 11.40 8.457e-10 96-
217	PR00381	KINESIN LIGHT CHAIN SIGNATURE	PR00381A 9.55 1.321e-10 360-
211	1100501		378
222	BL00514	Fibrinogen beta and gamma chains C-	BL00514C 17.41 2.358e-26 1166-
		terminal domain proteins.	1203 BL00514G 15.98 9.000e-15
			1289-1319 BL00514D 15.35
]			6.936e-12 1207-1220 BL00514F
			11.65 4.288e-10 1253-1268 BL00514H 14.95 8.636e-10 1318-
			1343
223	BL00325	Actin-depolymerizing proteins.	BL00325B 21.66 1.000e-40 93-
			139 BL00325A 24.83 9.333e-24
			61-93
224	BL00018	EF-hand calcium-binding domain	BL00018 7.41 1.450e-10 231-244
	protect	proteins.	DE01200D 10 50 1 502 10 57 50
225	PF01329	Pterin 4 alpha carbinolamine dhydratase.	PF01329B 18.52 1.692e-18 67-92 BL00211B 13.37 6.250e-18 1033-
228	BL00211	ABC transporters family proteins.	1065 BL00211B 13.37 6.250e-18 1033-
			2045-2077 BL00211A 12.23
}		1	1.900e-09 931-943
230	PR00761	BINDIN PRECURSOR SIGNATURE	PR00761A 5.81 9.366e-09 275-
			292
231	PR00049	WILM'S TUMOUR PROTEIN	PR00049D 0.00 3.500e-10 54-69
		SIGNATURE	
232	BL00412	Neuromodulin (GAP-43) proteins.	BL00412D 16.54 1.978e-10 109-
.			160 BL00412D 16.54 4.122e-09
233	BL01210	Caveolins proteins.	133-184 BL01210B 13.92 8.129e-09 106-
233	D£01210	Caveoims protems.	BL01210B 13.92 8.129e-09 106-
236	BL00939	Ribosomal protein L1e proteins.	BL00939F 17.27 5.393e-09 861-
		p.com Dropomo.	891
238	BL01252	Endogenous opioids neuropeptides	BL01252D 18.25 3.571e-28 205-
<u> </u>		precursors proteins.	233 BL01252B 19.09 5.034e-27

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SEQ ID NO:	ACCESSION NO.	DESCRIPTION	RESULTS*
110.			37-67 BL01252C 18.10 1.621e-21 164-190 BL01252A 14.22 7.107e- 18 14-34
239	BL00302	Eukaryotic initiation factor 5A hypusine proteins.	BL00302 14.81 1.000e-40 25-79
240	PR00420	AROMATIC-RING HYDROXYLASE (FLAVOPROTEIN MONOOXYGENASE) SIGNATURE	PR00420A 14.78 8.851e-13 26-49
241	PD02929	ADHESION GLYCOPROTEIN PRECURSOR I.	PD02929A 28.27 4.529e-09 235- 289
243	PD01066	PROTEIN ZINC FINGER ZINC- FINGER METAL-BINDING NU.	PD01066 19.43 8.527e-25 11-50
244	BL01270	Band 7 protein family proteins.	BL01270C 16.91 6.745e-17 115- 144 BL01270B 18.74 6.857e-17 76-115 BL01270E 13.03 6.016e- 15 182-211 BL01270D 20.87 9.160e-13 144-182
245	PF00791	Domain present in ZO-1 and Unc5-like netrin receptors.	PF00791B 28.49 6.305e-12 253- 308 PF00791B 28.49 1.909e-11 427-482 PF00791B 28.49 2.651e- 09 179-234 PF00791B 28.49 3.890e-09 112-167
246	PD00066	PROTEIN ZINC-FINGER METAL- BINDI.	PD00066 13.92 2.500e-13 277-290 PD00066 13.92 9.143e-12 193-206 PD00066 13.92 5.304e-11 165-178 PD00066 13.92 6.478e-11 249-262 PD00066 13.92 3.423e-10 221-234
247	BL00406	Actins proteins.	BL00406D 12.58 6.400e-20 465- 520 BL00406B 5.47 4.857e-14 249-304 BL00406E 8.44 1.000e- 11 522-572 BL00406C 6.75 5.449e-11 313-368
248	BL00951	ER lumen protein retaining receptor proteins.	BL00951C 19.35 1.000e-40 112- 161 BL00951A 15.10 7.750e-39 21-57 BL00951D 13.94 6.000e-38 161-196 BL00951B 14.23 3.100e- 31 57-88
252	BL01113	C1q domain proteins.	BL01113A 17.99 9.129e-15 200- 227 BL01113A 17.99 4.818e-14 194-221 BL01113A 17.99 7.818e- 14 182-209 BL01113A 17.99 1.730e-13 185-212 BL01113A 17.99 6.595e-13 191-218 BL01113A 17.99 6.077e-12 203- 230 BL01113A 17.99 9.182e-11 179-206 BL01113A 17.99 2.532e- 10 176-203 BL01113A 17.99 9.043e-10 218-245 BL01113A 17.99 9.426e-10 209-236 BL01113A 17.99 4.115e-09 137- 164
257	BL00845	CAP-Gly domain proteins.	BL00845 16.43 1.837e-21 466-491
259	PR00248	METABOTROPIC GLUTAMATE GPCR SIGNATURE	PR00248G 12.67 2.688e-09 53-78
260	BL00678	Trp-Asp (WD) repeat proteins proteins.	BL00678 9.67 3.400e-10 441-452 BL00678 9.67 5.800e-10 481-492 BL00678 9.67 8.800e-10 358-369
261	BL00678	Trp-Asp (WD) repeat proteins proteins.	BL00678 9.67 3.400e-10 415-426 BL00678 9.67 5.800e-10 455-466

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SEQ ID NO:	ACCESSION NO.	DESCRIPTION	RESULTS*
			BL00678 9.67 8.800e-10 332-343
262	BL00678	Trp-Asp (WD) repeat proteins proteins.	BL00678 9.67 3.400e-10 468-479 BL00678 9.67 5.800e-10 508-519 BL00678 9.67 8.800e-10 385-396
263	BL50002	Src homology 3 (SH3) domain proteins profile.	BL50002B 15.18 2.200e-10 415- 429
264	BL00049	Ribosomal protein L14 proteins.	BL00049C 17.38 3.040e-12 94- 130
265	PD01469	GLYCOPROTEIN PROTEIN PRECURSOR SA.	PD01469 20.59 2.091e-14 438-470
266	PD01469	GLYCOPROTEIN PROTEIN PRECURSOR SA.	PD01469 20.59 2.091e-14 279-311
267	BL00567	Phosphoribulokinase proteins.	BL00567A 10.66 1.161e-12 36-55
269	BL00049	Ribosomal protein L14 proteins.	BL00049C 17.38 2.688e-28 92- 128 BL00049B 18.42 6.806e-24 54-86 BL00049A 13.86 8.333e-19 19-42 BL00049D 13.47 5.765e-12 129-140
272	BL01115	GTP-binding nuclear protein ran proteins.	BL01115A 10.22 9.735e-12 14-58
273	PR00021	SMALL PROLINE-RICH PROTEIN SIGNATURE	PR00021A 4.31 1.911e-09 819- 832
275	PR00179	LIPOCALIN SIGNATURE	PR00179B 9.56 2.895e-13 124- 137 PR00179A 13.78 3.250e-11 36-49 PR00179C 19.02 6.040e-11 154-170
276	PR00449	TRANSFORMING PROTEIN P21 RAS SIGNATURE	PR00449A 13.20 8.364e-17 22-44 PR00449C 17.27 1.000e-13 62-85 PR00449E 13.50 4.000e-12 172- 195 PR00449B 14.34 5.680e-10 45-62
277	BL00140	Ubiquitin carboxyl-terminal hydrolase family I cysteine activ.	BL00140D 22.64 1.000e-40 161- 205 BL00140C 11.80 9.053e-30 79-104 BL00140A 15.96 9.400e- 28 5-35 BL00140B 12.29 4.649e- 17 37-55
278	PD02712	ELEMENT TRANSPOSASE FOR TRANSPOSON TRANSPOSABLE.	PD02712A 23.03 8.013e-09 47-83
279	BL00678	Trp-Asp (WD) repeat proteins proteins.	BL00678 9.67 1.474e-09 100-111
282	DM00892	3 RETROVIRAL PROTEINASE.	DM00892C 23.55 4.767e-21 864- 898
283	BL00048	Protamine P1 proteins.	BL00048 6.39 9.550e-09 56-83
286	PR00081	GLUCOSE/RIBITOL DEHYDROGENASE FAMILY SIGNATURE	PR00081A 10.53 1.878e-11 36-54
287	PR00310	ANTI-PROLIFERATIVE PROTEIN BTG1 FAMILY SIGNATURE	PR00310B 10.59 4.231e-17 29-59 PR00310D 9.10 6.679e-16 89-119
289	PD01066	PROTEIN ZINC FINGER ZINC- FINGER METAL-BINDING NU.	PD01066 19.43 7.000e-36 37-76
293	BL00979	G-protein coupled receptors family 3 proteins.	BL00979L 20.63 3.800e-12 111- 152
295	PD02411	PROTEIN TRANSCRIPTION REGULATION NUCLEAR.	PD02411 21.89 7.000e-16 195-229
296	BL01064	Pyridoxamine 5'-phosphate oxidase proteins.	BL01064A 27.84 8.313e-28 77- 129 BL01064C 15.22 7.136e-25 202-235
297	BL00030	Eukaryotic RNA-binding region RNP-1 proteins.	BL00030A 14.39 2.929e-13 37-56 BL00030B 7.03 1.900e-11 167- 177 BL00030A 14.39 2.000e-10 128-147

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CTPO	ACCESSION	DESCRIPTION	RESULTS*
SEQ		DESCRIPTION	KESOLIS.
ID ID	NO.	•	
NO:	DI 01102	ubiE/COQ5 methyltransferase family	BL01183B 21.31 6.660e-12 143-
298	BL01183	proteins.	188
000	DI 01000	1 •	BL01279A 24.27 5.862e-11 57-
299	BL01279	Protein-L-isoaspartate(D-aspartate) O-	105
	DI 00101	methyltransferase signa. Cytochrome b5 family, heme-binding	BL00191K 17.38 4.951e-27 184-
301	BL00191		228 BL00191J 11.37 6.447e-17
		domain proteins.	128-150
	72.60000	A DECEMBAL DE OCCUPIA DE	DM00892C 23.55 3.893e-16 33-67
302	DM00892	3 RETROVIRAL PROTEINASE.	
306	PF01140	Matrix protein (MA), p15.	PF01140D 15.54 2.988e-09 416-
			451
307	PR00245	OLFACTORY RECEPTOR	PR00245A 18.03 4.818e-21 59-81
1		SIGNATURE	PR00245C 7.84 5.154e-20 238-
			254 PR00245D 10.47 4.000e-15
			274-286 PR00245B 10.38 8.200e-
			15 177-192 PR00245E 12.40
			5.714e-12 291-306
309	BL00203	Vertebrate metallothioneins proteins.	BL00203 13.94 2.245e-10 612-658
310	BL00237	G-protein coupled receptors proteins.	BL00237A 27.68 7.632e-23 119-
			159 BL00237C 13.19 3.864e-15
1 1			251-278 BL00237D 11.23 3.739e-
			12 312-329
311	BL00380	Rhodanese proteins.	BL00380D 15.90 8.200e-28 110-
			136 BL00380G 11.26 5.800e-16
			267-280 BL00380B 14.77 7.000e-
		·	14 49-62 BL00380F 9.76 5.886e-
			13 203-214 BL00380C 15.67
1	lii		7.387e-13 82-98 BL00380E 12.44
			7.000e-11 181-193 BL00380A
		•	10.48 1.000e-09 10-20
312	BL00227	Tubulin subunits alpha, beta, and gamma	BL00227B 19.29 1.000e-40 50-
		proteins.	105 BL00227C 25.48 1.000e-40
			111-163 BL00227D 18.46 1.000e-
			40 220-274 BL00227F 21.16
			1.000e-40 372-426 BL00227A
			24.55 3.250e-39 1-35 BL00227E
			24.15 8.500e-34 324-359
327	BL00232	Cadherins extracellular repeat proteins	BL00232B 32.79.7.362e-21 225-
		domain proteins.	273 BL00232B 32.79 2.588e-17
			435-483 BL00232B 32.79 6.301e-
			15 116-164 BL00232B 32.79
]			6.769e-13 330-378 BL00232C
			10.65 9.341e-12 223-241
			BL00232C 10.65 5.696e-11 328-
]			346 BL00232C 10.65 3.942e-10
			433-451
329	PD02749	TRANSCRIPTION PROTEIN FACTOR	PD02749B 12.75 2.241e-37 35-71
		BTF3 REGULATION NUCL.	PD02749C 13.96 4.892e-28 87-
1			121 PD02749A 9.56 6.000e-15 2-
			15
330	PR00391	PHOSPHATIDYLINOSITOL	PR00391E 12.50 7.785e-15 211-
		TRANSFER PROTEIN SIGNATURE	231 PR00391B 8.39 1.000e-13
1			83-104 PR00391D 12.21 9.328e-
		{	13 191-207 PR00391A 7.83
			5.390e-11 16-36
332	BL01030	RNA polymerases M / 15 Kd subunits	BL01030 23.44 1.818e-23 87-125
		proteins.	
337	PD01066	PROTEIN ZINC FINGER ZINC-	PD01066 19.43 2.929e-32 6-45
	1201000	FINGER METAL-BINDING NU.	
340	PD02711	SYNTHASE	PD02711B 14.26 1.973e-20 944-

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SEQ ID NO:	ACCESSION NO.	DESCRIPTION	RESULTS*
		PHOSPHORIBOSYLFORMYLGLY.	968
343	BL00223	Annexins repeat proteins domain proteins.	BL00223C 24.79 1.000e-40 245- 300 BL00223B 28.47 8.714e-38 168-218 BL00223A 15.59 8.250e- 27 98-132 BL00223A 15.59 8.750e-27 26-60 BL00223C 24.79 9.438e-16 13-68 BL00223C 24.79 2.735e-15 85-140 BL00223A 15.59 2.253e-11 258-292
346	PR00345	STATHMIN FAMILY SIGNATURE	PR00345B 7.12 2.800e-28 81-110 PR00345E 8.54 7.652e-28 158- 183 PR00345C 4.54 9.100e-28 110-134 PR00345D 10.97 1.964e- 24 134-158 PR00345A 13.46 5.645e-16 52-71
347	BL00586	Ribosomal protein L16 proteins.	BL00586B 17.00 3.215e-15 184- 221
348	PR00388	3',5'-CYCLIC NUCLEOTIDE CLASS II PHOSPHODIESTERASE SIGNATURE	PR00388A 10.45 2.778e-09 86- 105
351	BL00018	EF-hand calcium-binding domain proteins.	BL00018 7.41 3.118e-11 160-173 BL00018 7.41 2.350e-10 244-257
354	BL00678	Trp-Asp (WD) repeat proteins proteins.	BL00678 9.67 1.947e-09 256-267
358	DM01206	CORONAVIRUS NUCLEOCAPSID PROTEIN.	DM01206B 10.69 3.278e-09 175- 195 DM01206B 10.69 6.696e-09 183-203 DM01206B 10.69 8.633e-09 132-152 DM01206B 10.69 8.861e-09 181-201 DM01206B 10.69 9.316e-09 177- 197
361	PD01498	OXIDASE BIOSYNTHESIS OXIDOREDUCTASE PORP.	PD01498C 24.90 6.880e-14 219- 263
362	PD01498	OXIDASE BIOSYNTHESIS OXIDOREDUCTASE PORP.	PD01498C 24.90 6.880e-14 219- 263
365	BL00178	Aminoacyl-transfer RNA synthetases class-I proteins.	BL00178B 7.11 1.000e-11 589- 600 BL00178A 14.23 8.500e-09 46-56
366	BL00523	Sulfatases proteins.	BL00523E 19.27 1.000e-23 318- 348 BL00523A 13.36 5.500e-16 30-47 BL00523B 8.64 1.964e-13 78-90 BL00523C 12.64 9.625e-13 129-140 BL00523G 9.46 5.500e- 10 506-516
369	BL00107	Protein kinases ATP-binding region proteins.	BL00107A 18.39 4.818e-09 21-52
370	BL00880	Acyl-CoA-binding protein.	BL00880 17.52 1.000e-40 75-125
371	BL00107	Protein kinases ATP-binding region proteins.	BL00107A 18.39 1.000e-23 276- 307 BL00107B 13.31 1.692e-12 342-358
372	PR00211	GLUTELIN SIGNATURE .	PR00211B 0.86 6.602e-11 326- 347 PR00211B 0.86 6.106e-10 320-341 PR00211B 0.86 3.167e- 09 333-354
373	BL00279	Membrane attack complex components / perforin proteins.	BL00279E 37.11 9.349e-10 749- 7.97
375	PD01066	PROTEIN ZINC FINGER ZINC- FINGER METAL-BINDING NU.	PD01066 19.43 1.231e-33 10-49
377	- PD01066	PROTEIN ZINC FINGER ZINC- FINGER METAL-BINDING NU.	PD01066 19.43 7.563e-28 10-49
379	BL00598	Chromo domain proteins.	BL00598 14.45 5.781e-16 3-25

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SEQ	ACCESSION	DESCRIPTION	RESULTS*
ID	NO.		
NO:			
380	PR00413	HALOACID	PR00413D 11.28 8.941e-09 864-
		DEHALOGENASE/EPOXIDE	878
		HYDROLASE FAMILY SIGNATURE	
383	PR00413	HALOACID	PR00413D 11.28 8.941e-09 864-
		DEHALOGENASE/EPOXIDE	878
		HYDROLASE FAMILY SIGNATURE	
387	BL01060	Flagella transport protein fliP family	BL01060A 15.65 1.535e-09 131-
		proteins.	174
388	PR00209	ALPHA/BETA GLIADIN FAMILY	PR00209B 4.88 6.318e-11 1009-
		SIGNATURE	1028
389	PR00837	ALLERGEN V5/TPX-1 FAMILY	PR00837B 11.64 1.000e-10 469-
		SIGNATURE	483
391	BL00240	Receptor tyrosine kinase class III	BL00240B 24.70 7.907e-10 118-
	7700014	proteins.	142
392	PR00014	FIBRONECTIN TYPE III REPEAT	PR00014D 12.04 8.412e-10 691-
	DD 00014	SIGNATURE	706 PR00014D 12.04 8.412e-10 706-
393	PR00014	FIBRONECTIN TYPE III REPEAT	721
204	DI 01200	SIGNATURE	BL01209 9.31 3.368e-15 47-60
394	BL01209	LDL-receptor class A (LDLRA) domain proteins.	BL01209 9.31 5.500e-13 47-60 BL01209 9.31 5.500e-13 92-105
395	BL00634	Ribosomal protein L30 proteins.	BL00634 34.38 4.090e-13 70-121
393	BL01013	Oxysterol-binding protein family	BL01013D 26.81 8.000e-26 358-
390	DL01013	proteins.	402 BL01013A 25.14 7.231e-21
-		protents.	45-81 BL01013C 9.97 1.000e-13
		·	132-142 BL01013B 11.33 1.000e-
			11 110-121
397	BL00930	Peripherin / rom-1 proteins.	BL00930E 17.80 1.000e-40 56-92
]		1 or proteins	BL00930D 9.12 4.632e-37 12-56
			BL00930F 16.91 2.800e-36 92-
			133
400	PR00780	LEUSERPIN 2 SIGNATURE	PR00780B 4.89 4.491e-09 262-
			285
401	PR00819	CBXX/CFQX SUPERFAMILY	PR00819B 10.83 7.158e-11 4-20
·		SIGNATURE	
403	BL00381	Endopeptidase Clp serine proteins.	BL00381C 23.84 1.250e-32 150-
			194 BL00381A 16.48 2.286e-22
			74-111 BL00381B 21.42 8.326e-
			14 78-130
405	BL01105	Ribosomal protein L35Ae proteins.	BL01105A 17.37 1.000e-40 4-49
			BL01105B 12.95 1.000e-40 68-
406	DY 00244	CATA	108
406	BL00344	GATA-type zinc finger domain proteins.	BL00344 17.99 7.000e-12 814-852 PR00211B 0.86 9.750e-09 73-94
407	PR00211	GLUTELIN SIGNATURE	PR00211B 0.86 9.750e-09 73-94 PR00910A 2.51 4.321e-09 9-22
409	PR00910	LUTEOVIRUS ORF6 PROTEIN SIGNATURE	FRUUTIUM 2.31 4.3216-09 9-22
410	BL00762	WHEP-TRS domain proteins.	BL00762A 23.43 1.000e-28 752-
410	DLUU/02	WITEF-IRS domain proteins.	789 BL00762A 23.43 1.000e-28 732-
			903-940 BL00762A 23.43 5.415e-
			18 825-862 BL00762B 16.14
			8.759e-12 1154-1168
412	BL00690	DEAH-box subfamily ATP-dependent	BL00690B 13.38 5.320e-15 262-
112	2200070	helicases proteins.	280 BL00690A 6.87 1.818e-13
1		P. C. C. C. C. C. C. C. C. C. C. C. C. C.	230-240
415	BL00227	Tubulin subunits alpha, beta, and gamma	BL00227B 19.29 1.000e-40 52-
		proteins.	107 BL00227C 25.48 1.000e-40
		•	113-165 BL00227D 18.46 1.000e-
			40 222-276 BL00227F 21.16
			1.000e-40 382-436 BL00227E
		<u> </u>	24.15 1.750e-34 326-361

SEQ	ACCESSION	DESCRIPTION	RESULTS*
ID	NO.		
NO:			
			BL00227A 24.55 1.000e-33 1-35
416	PF00992	Troponin.	PF00992A 16.67 1.711e-09 557-
		•	592
418	BL00541	Nuclear transition protein 1 proteins.	BL00541 8.44 9.875e-09 256-310
419	BL00541	Nuclear transition protein 1 proteins.	BL00541 8.44 9.875e-09 197-251
420	PF00856	SET domain proteins.	PF00856A 26.14 9.074e-13 901-
		_	938 PF00856B 16.42 2.397e-12
			951-973
421	BL00678	Trp-Asp (WD) repeat proteins proteins.	BL00678 9.67 8.200e-12 33-44
423	PD01066	PROTEIN ZINC FINGER ZINC-	PD01066 19.43 8.600e-30 130-169
		FINGER METAL-BINDING NU.	
424	PF00564	Octicosapeptide repeat proteins.	PF00564B 24.74 1.305e-17 421-
			472
426	PR00988	URIDINE KINASE SIGNATURE	PR00988A 6.39 4.569e-12 3-21
427	PR00988	URIDINE KINASE SIGNATURE	PR00988A 6.39 4.569e-12 3-21
428	BL00478	LIM domain proteins.	BL00478B 14.79 3.250e-13 115-
			130 BL00478B 14.79 9.036e-13
			50-65
431	BL00282	Kazal serine protease inhibitors family	BL00282 16.88 8.875e-12 464-487
		proteins.	DD 00000D 00 50 5 500
432	PD00930	PROTEIN GTPASE DOMAIN	PD00930B 33.72 7.800e-18 316-
		ACTIVATION.	357 PD00930A 25.62 9.617e-12
			125-151 PD00930B 33.72 2.521e-
			10 214-255
433	PD01066	PROTEIN ZINC FINGER ZINC-	PD01066 19.43 4.649e-34 34-73
		FINGER METAL-BINDING NU.	PR00449A 13.20 7.563e-11 56-78
434	PR00449	TRANSFORMING PROTEIN P21 RAS	PR00449A 13.20 7.303e-11 30-78
100	77700100	SIGNATURE H+-TRANSPORTING ATPASE	PR00120C 9.90 5.800e-19 705-
436	PR00120	(PROTON PUMP) SIGNATURE	722.
427	BL00115	Eukaryotic RNA polymerase II	BL00115T 8.45 7.273e-29 1208-
437	BL00113	heptapeptide repeat proteins.	1242 BL00115Q 18.08 2.776e-21
		neptapeptide repeat protonis.	953-983 BL00115Y 11.86 8.000e-
			17 1604-1650 BL00115M 19.19
			8.130e-16 731-774 BL00115H
		•	14.34 9.392e-16 463-496
			BL00115A 15.44 7.414e-15 43-82
			BL00115R 6.50 6.128e-14 983-
			1010 BL00115J 16.71 9.289e-14
			591-617 BL00115I 8.33 4.336e-
			13 535-590 BL00115L 12.25
			5.939e-13 662-694 BL00115G
			11.65 6.011e-13 435-463
			BL00115K 15.03 3.417e-10 617-
			659 BL00115O 16.76 5.805e-10
			863-913 BL00115P 11.54 7.538e-
			10 913-953 BL00115S 18.24
	*		7.968e-10 1010-1052 BL00115U
		7110 6	10.34 4.475e-09 1242-1265
438	PF00628	PHD-finger.	PF00628 15.84 4.536e-10 219-234
440	PD01066	PROTEIN ZINC FINGER ZINC-	PD01066 19.43 6.351e-34 10-49
	DDOOCO	FINGER METAL-BINDING NU.	PR00309A 9.68 5.250e-24 32-55
441	PR00309	ARRESTIN SIGNATURE	PR00309A 9.68 5.250e-24 32-35 PR00309D 7.09 4.938e-23 290-
			309 PR00309B 7.81 2.800e-21
			69-88 PR00309C 8.22 1.621e-19
			165-183 PR00309E 9.82 9.438e-
1			15 374-389
442	BL00600	Aminotransferases class-III pyridoxal-	BL00600B 19.60 7.324e-14 103-
742	DE00000	1 minion minion mono orang-rit blingorm.	122300002 27.00 7.00.00 7.00

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SEQ ID NO:	ACCESSION NO.	DESCRIPTION	RESULTS*
		phosphate attachment si.	129 BL00600G 12.43 2.125e-12 306-325 BL00600F 8.77 8.105e- 12 271-284 BL00600E 16.43 3.167e-11 228-257 BL00600D 8.71 8.650e-09 207-221
443	BL00972	Ubiquitin carboxyl-terminal hydrolases family 2 proteins.	BL00972A 11.93 3.160e-18 69-87
444	BL00349	CTF/NF-I proteins.	BL00349A 10.07 1.000e-40 8-54 BL00349C 9.33 1.000e-40 82-125 BL00349E 10.79 1.000e-40 152- 195 BL00349F 11.81 1.000e-40 213-255 BL00349H 15.70 7.387e- 36 361-399 BL00349B 10.51 2.227e-34 54-82 BL00349D 11.70 9.100e-34 125-152 BL00349G 19.72 5.781e-30 323-356
445	BL00154	E1-E2 ATPases phosphorylation site proteins.	BL00154F 8.23 8.941e-21 271- 295 BL00154E 20.37 2.620e-15 124-165
448	DM00215	PROLINE-RICH PROTEIN 3.	DM00215 19.43 4.882e-11 82-115 DM00215 19.43 6.492e-09 87-120
451	BL01283	T-box domain proteins.	BL01283A 24.15 3.100e-40 112- 160 BL01283D 11.70 6.000e-39 253-286 BL01283B 23.17 6.538e- 38 170-212 BL01283C 13.05 7.750e-19 222-236
452	PR00420	AROMATIC-RING HYDROXYLASE (FLAVOPROTEIN MONOOXYGENASE) SIGNATURE	PR00420A 14.78 2.579e-11 3-26
453	PR00162	RIESKE 2FE-2S SUBUNIT SIGNATURE	PR00162B 12.77 7.429e-17 215- 228 PR00162A 9.35 2.324e-14 193-205 PR00162C 8.10 7.120e- 14 227-240
454	PD01066	PROTEIN ZINC FINGER ZINC- FINGER METAL-BINDING NU.	PD01066 19.43 7.000e-30 87-126
456	BL00027	'Homeobox' domain proteins.	BL00027 26.43 9.333e-18 1149- 1192
457	PD01066	PROTEIN ZINC FINGER ZINC- FINGER METAL-BINDING NU.	PD01066 19.43 2.737e-24 16-55
459	BL00290	Immunoglobulins and major histocompatibility complex proteins.	BL00290A 20.89 1.529e-14 154- 177 BL00290B 13.17 9.000e-12 214-232
460	PR00413	HALOACID DEHALOGENASE/EPOXIDE HYDROLASE FAMILY SIGNATURE	PR00413F 14.91 7.333e-11 193- 214 PR00413E 15.78 5.714e-09 175-192
463	PR00759	BASIC PROTEASE (KUNITZ-TYPE) INHIBITOR FAMILY SIGNATURE	PR00759B 11.26 8.385e-09 74-85
466	BL00019	Actinin-type actin-binding domain proteins.	BL00019D 15.33 4.200e-19 300- 330
467	BL00019	Actinin-type actin-binding domain proteins.	BL00019D 15.33 4.200e-19 300- 330
469	PR00153	CYCLOPHILIN PEPTIDYL-PROLYL CIS-TRANS ISOMERASE SIGNATURE	PR00153D 11.99 3.250e-15 510- 523 PR00153C 11.01 4.682e-14 495-511 PR00153E 9.10 8.548e- 14 523-539 PR00153B 11.57 1.720e-13 452-465
470	BL00491	Aminopeptidase P and proline dipeptidase proteins.	BL00491C 12.15 3.912e-09 557- 572
471	PD00289	PROTEIN SH3 DOMAIN REPEAT	PD00289 9.97 1.000e-14 1482-

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SEQ	ACCESSION	DESCRIPTION	RESULTS*
ID ID	NO.	DESCRIPTION	100010
NO:	110.		
140.		PRESYNA.	1496 PD00289 9.97 8.650e-11
		TRESTITE.	1122-1136
474	BL50040	Elongation factor 1 gamma chain profile.	BL50040D 17.41 1.000e-40 279-
4/4	DL30040	Diongation ratio 1 gamma onum provinc.	329 BL50040E 18.79 1.000e-40
			333-388 BL50040F 18.99 5.320e-
			40 390-428 BL50040C 22.62
			3,739e-38 141-184 BL50040B
			13.65 7.000e-30 59-85 BL50040A
			12.98 1.450e-14 10-22
475	BL01144	Ribosomal protein L31e proteins.	BL01144 25.07 1.000e-40 22-74
476	PR00007	COMPLEMENT CIQ DOMAIN	PR00007C 15.60 2.421e-21 589-
.,,	1100007	SIGNATURE	611 PR00007B 14.16 3.500e-21
			544-564 PR00007A 19.33 6.897e-
			20 517-544 PR00007D 9.64
			6.571e-12 623-634
477	BL50002	Src homology 3 (SH3) domain proteins	BL50002A 14.19 5.846e-10 170-
		profile.	189
479	DM01970	0 kw ZK632.12 YDR313C	DM01970B 8.60 9.500e-17 967-
		ENDOSOMAL III.	980
480	PR00868	DNA-POLYMERASE FAMILY A (POL	PR00868C 13.76 5.688e-17 284-
		I) SIGNATURE	308 PR00868A 16.33 3.186e-13
		•	224-247 PR00868H 12.51 3.388e-
			13 431-448 PR00868I 10.87
			7.938e-11 462-476 PR00868E
			13.19 1.608e-10 340-366
481	BL00027	'Homeobox' domain proteins.	BL00027 26.43 9.182e-22 53-96
482	BL00061	Short-chain dehydrogenases/reductases	BL00061B 25.79 3.647e-21 188-
		family proteins.	226
483	BL50002	Src homology 3 (SH3) domain proteins	BL50002A 14.19 1.750e-12 1032-
		profile.	1051
485	PF00023	Ank repeat proteins.	PF00023A 16.03 9.625e-10 760-
			776 PF00023A 16.03 3.571e-09 715-731
406	DD00070	RECEPTOR INTERLEUKIN-1	PD02870B 18.83 9.262e-20 103-
486	PD02870	PRECURSOR.	136 PD02870D 15.74 9.426e-09
		PRECORSOR.	201-236
487	PR00370	FLAVIN-CONTAINING	PR00370G 10.45 3.769e-28 471-
407	PR00370	MONOOXYGENASE (FMO)	493 PR00370B 10.91 1.000e-24
		SIGNATURE .	27-46 PR00370C 12.72 4.000e-21
		SIGNATIONS .	140-157 PR00370E 11.96 9.229e-
			21 320-339 PR00370D 16.33
			1.750e-20 185-204 PR00370F
			17.75 7.395e-20 375-395
			PR00370A 3.35 2.038e-18 4-20
489	PD01675	GLYCOPROTEIN MAJOR ENVELOPE	PD01675C 19.89 2.330e-10 55-89
		PROBABLE U3.	
492	BL00211	ABC transporters family proteins.	BL00211A 12.23 5.050e-09 45-57
493	BL00211	ABC transporters family proteins.	BL00211A 12.23 5.050e-09 45-57
494	BL00211	ABC transporters family proteins.	BL00211A 12.23 5.050e-09 58-70
495	BL00027	'Homeobox' domain proteins.	BL00027 26.43 6.786e-12 509-552
			BL00027 26.43 9.143e-12 319-362
			BL00027 26.43 2.600e-11 627-670
			BL00027 26.43 3.625e-10 779-822
497	BL00107	Protein kinases ATP-binding region	BL00107A 18.39 5.800e-22 214-
		proteins.	245 BL00107B 13.31 1.000e-13
			281-297 BL00107A 18.39 3.520e-
J			
			13 583-614 BL00107B 13.31
499	BL00383	Tyrosine specific protein phosphatases	13 583-614 BL00107B 13.31 8.615e-12 652-668 BL00383E 10.35 1.000e-14 1902-

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SEQ ID NO:	ACCESSION NO.	DESCRIPTION	RESULTS*
NO:		proteins.	1913 BL00383D 11.92 3.077e-14 1862-1875 BL00383A 13.34 5.500e-14 1730-1745 BL00383C 10.10 2.000e-13 1785-1796 BL00383F 15.51 9.069e-12 1940- 1956 BL00383B 7.61 1.692e-11 1755-1764
501	PR00019	LEUCINE-RICH REPEAT SIGNATURE	PR00019B 11.36 1.360e-09 136- 150 PR00019A 11.19 1.667e-09 91-105 PR00019B 11.36 4.600e- 09 160-174
503	BL00226	Intermediate filaments proteins.	BL00226D 19.10 1.000e-40 367- 414 BL00226B 23.86 6.143e-27 195-243 BL00226A 12.77 7.840e- 14 96-111 BL00226C 13.23 2.600e-13 309-340 BL00226C 13.23 6.143e-12 266-297 BL00226B 23.86 1.209e-09 146- 194
505	PD02407	3-BISPHOSPHOGLYCERATE- INDEPENDENT PHOSPHOGLYCER.	PD02407F 7.61 6.739e-09 916- 930
506	PF00632	HECT-domain (ubiquitin-transferase).	PF00632C 20.66 9.830e-19 991- 1023 PF00632B 18.45 1.155e-11 940-968
507	BL01082	Ribosomal protein L7Ae proteins.	BL01082 20.37 4.273e-20 76-116
508	BL00678	Trp-Asp (WD) repeat proteins proteins.	BL00678 9.67 2.421e-09 493-504
509	BL00678	Trp-Asp (WD) repeat proteins proteins.	BL00678 9.67 2.421e-09 473-484
510	PR00320	G-PROTEIN BETA WD-40 REPEAT SIGNATURE	PR00320B 12.19 4.774e-11 567-582 PR00320B 12.19 5.886e-10 763-778 PR00320C 13.01 6.760e-10 567-582 PR00320A 16.74 7.618e-10 846-861 PR00320A 16.74 3.415e-09 763-778 PR00320A 16.74 6.268e-09 567-582
511	BL00479	Phorbol esters / diacylglycerol binding domain proteins.	BL00479C 12.01 3.250e-12 170- 183
512	BL50058	G-protein gamma subunit profile.	BL50058 27.23 7.494e-09 10-58
513	BL00524	Somatomedin B domain proteins.	BL00524A 9.65 8.925e-14 80-101
515	BL00041	Bacterial regulatory proteins, araC family proteins.	BL00041 23.99 1.964e-19 492-524
516	PD00066	PROTEIN ZINC-FINGER METAL- BINDI.	PD00066 13.92 8.500e-13 391-404
517	BL00415	Synapsins proteins.	BL00415E 4.82 9.291e-09 959- 996
518	PR00109	TYROSINE KINASE CATALYTIC DOMAIN SIGNATURE	PR00109B 12.27 9.471e-12 126- 145
519	BL00290	Immunoglobulins and major histocompatibility complex proteins.	BL00290B 13.17 4.750e-09 47-65
522	PR00505	D12 CLASS N6 ADENINE-SPECIFIC DNA METHYLTRANSFERASE SIGNATURE	PR00505A 14.15 7.128e-09 364- 381
525	BL00312	Glycophorin A proteins.	BL00312B 9.22 5.781e-10 891- 920
528	PD01066	PROTEIN ZINC FINGER ZINC- FINGER METAL-BINDING NU.	PD01066 19.43 2.500e-32 16-55
529	PR00254	NICOTINIC ACETYLCHOLINE RECEPTOR SIGNATURE	PR00254D 15.50 4.000e-17 131- 150 PR00254A 11.23 4.706e-14 61-78 PR00254C 11.36 4.000e-12

SEQ ID NO:	ACCESSION NO.	DESCRIPTION	RESULTS*
			113-126 PR00254B 12.97 1.486e- 11 95-110
531	BL00741	Guanine-nucleotide dissociation stimulators CDC24 family sign.	BL00741B 14.27 6.870e-16 787- 810
532	PR00193	MYOSIN HEAVY CHAIN SIGNATURE	PR00193D 14.36 3.143e-34 447- 476 PR00193C 12.60 7.632e-32 216-244 PR00193B 11.69 7.750e- 29 167-193 PR00193A 15.41 2.588e-22 111-131 PR00193E 19.47 2.200e-21 501-530
533	PD02870	RECEPTOR INTERLEUKIN-1 PRECURSOR.	PD02870B 18.83 5.596e-09 348- 381
535	PR00683	SPECTRIN PLECKSTRIN HOMOLOGY DOMAIN SIGNATURE	PR00683D 15.87 2.452e-10 465- 484
536	BL00027	'Homeobox' domain proteins.	BL00027 26.43 6.684e-24 164-207
538	PR00239	MOLLUSCAN RHODOPSIN C- TERMINAL TAIL SIGNATURE	PR00239E 1.58 2.739e-09 225- 237
539	BL00406	Actins proteins.	BL00406C 6.75 1.000e-40 157- 212 BL00406B 5.47 6.143e-37 90-145 BL00406D 12.58 4.600e- 36 291-346 BL00406E 8.44 2.200e-33 364-414 BL00406A 9.95 4.441e-23 7-42
540	PR00456	RIBOSOMAL PROTEIN P2 SIGNATURE	PR00456E 3.06 9.625e-10 44-59
541	PR00456	RIBOSOMAL PROTEIN P2 SIGNATURE	PR00456E 3.06 9.625e-10 44-59
542	PF00023	Ank repeat proteins.	PF00023A 16.03 7.857e-11 138- 154
544	PF00642	Zinc finger C-x8-C-x5-C-x3-H type (and similar).	PF00642 11.59 9.082e-10 838-849
546	BL00383	Tyrosine specific protein phosphatases proteins.	BL00383E 10.35 4.115e-10 104- 115
547	BL01226	Hydroxymethylglutaryl-coenzyme A synthase proteins.	BL01226A 13.79 1.000e-40 50-89 BL01226C 13.51 1.000e-40 127- 167 BL01226D 11.60 1.000e-40 174-210 BL01226E 13.74 1.000e- 40 212-253 BL01226H 17.74 1.000e-40 386-434 BL01226I 25.06 1.000e-40 460-508 BL01226G 15.76 3.483e-32 292- 321 BL01226B 13.35 1.818e-31 95-127 BL01226F 9.78 8.714e-23 253-271
549	BL00964	Syndecans proteins.	BL00964B 12.05 2.426e-10 1246- 1289
551	DM01930	2 kw FINGER SMCX SMCY YDR096W.	DM01930E 15.41 1.367e-37 170- 215 DM01930F 14.16 8.232e-28 267-303 DM01930B 19.86 9.163e-10 37-71
552	BL00195	Glutaredoxin proteins.	BL00195B 15.31 7.158e-09 9-29
554	BL00383	Tyrosine specific protein phosphatases proteins.	BL00383E 10.35 2.756e-12 436- 447
555	PR00403	WW DOMAIN SIGNATURE	PR00403B 12.19 7.612e-11 122- 137 PR00403A 16.82 3.912e-10 107-121 PR00403B 12.19 2.068e- 09 76-91
558	PR00380	KINESIN HEAVY CHAIN SIGNATURE	PR00380A 14.18 2.714e-26 76-98 PR00380D 9.93 3.000e-24 275-

SEQ	ACCESSION	DESCRIPTION	RESULTS*
IID	NO.		
NO:			
			297 PR00380C 13.18 5.154e-20
			226-245 PR00380B 12.64 9.400e-
			20 195-213
559 ·	BL00518	Zinc finger, C3HC4 type (RING finger),	BL00518 12.23 5.333e-09 522-531
		proteins.	
561	PD01795	PROTEIN AMINOPEPTIDASE	PD01795B 11.56 2.333e-12 159-
		PRECURSOR HYDROLASE SIGNA.	172 PD01795A 10.27 1.000e-09
			135-144
562	PD01795	PROTEIN AMINOPEPTIDASE	PD01795B 11.56 2.333e-12 110-
		PRECURSOR HYDROLASE SIGNA.	123 PD01795A 10.27 1.000e-09
			86-95
563	BL00018	EF-hand calcium-binding domain	BL00018 7.41 1.391e-09 41-54
		proteins.	
565	BL00348	p53 tumor antigen proteins.	BL00348F 23.19 4.143e-09 188-
505	22003.0	poo tamor amgan proteam.	231
567	PD00301	PROTEIN REPEAT MUSCLE	PD00301B 5.49 4.115e-09 284-
307	1 000301	CALCIUM-BI.	295
569	PF00850	Histone deacetylase family.	PF00850E 8.88 6.553e-21 756-782
309	1100000	Historie deacetylase lamily.	PF00850D 14.76 1.519e-16 722-
			746 PF00850F 15.70 1.118e-11
			794-827 PF00850G 22.75 8.375e-
			11 833-875
570	PD00289	PROTEIN SH3 DOMAIN REPEAT	PD00289 9.97 4.960e-10 137-151
3/0	PD00289	PRESYNA.	PD00289 9.97 4.9006-10 137-131
571	DI 00510		BL00518 12.23 8.800e-11 44-53
571	BL00518	Zinc finger, C3HC4 type (RING finger),	BL00318 12.23 8.800e-11 44-33
600	DI 00000	proteins.	BL00299 28.84 1.123e-11 123-175
573	BL00299	Ubiquitin domain proteins.	
574	PF01140	Matrix protein (MA), p15.	PF01140D 15.54 3.700e-10 986-
	=====		1021
576	BL00284	Serpins proteins.	BL00284C 28.56 5.200e-26 200-
			242 BL00284A 15.64 4.913e-18
			71-95 BL00284B 17.99 7.261e-15
			173-194 BL00284D 16.34 5.846e-
			13 306-333 BL00284E 19.15
	7701066		7.429e-12 387-412
579	PD01066	PROTEIN ZINC FINGER ZINC-	PD01066 19.43 6.553e-29 15-54
		FINGER METAL-BINDING NU.	
580	。 BL50001	Src homology 2 (SH2) domain proteins	BL50001B 17.40 4.500e-12 1010-
		profile.	1031
581	PD00930	PROTEIN GTPASE DOMAIN	PD00930B 33.72 3.189e-22 608-
		ACTIVATION.	649 PD00930A 25.62 6.806e-17
			505-531
584	BL00612	Osteonectin domain proteins.	BL00612B 11.35 2.034e-11 93-
			126
585	DM01551	kw OSTEOINDUCTIVE YOPM	DM01551C 14.62 8.859e-10 102-
		MEMBRANE OUTER.	122
586	PF00628	PHD-finger.	PF00628 15.84 3.455e-12 235-250
587	BL00027	'Homeobox' domain proteins.	BL00027 26.43 6.063e-10 85-128
588	PR00326	GTP1/OBG GTP-BINDING PROTEIN	PR00326A 8.75 7.525e-16 227-
		FAMILY SIGNATURE	248 PR00326C 9.79 6.760e-15
			276-292 PR00326D 19.09 6.657e-
			13 293-312 PR00326B 16.74
			9.229e-13 248-267
589	BL00422	Granins proteins.	BL00422A 28.34 7.429e-09 2349-
			2378
590	BL00415	Synapsins proteins.	BL00415N 4,29 9.794e-10 295-
		,	339
591	BL00128	Alpha-lactalbumin / lysozyme C proteins.	BL00128A 20.76 3.423e-13 35-65
		, proteins.	BL00128C 19.34 2.980e-11 110-
<u> </u>	<u> </u>	<u> </u>	

SEQ	ACCESSION	DESCRIPTION	RESULTS*
ID	NO.		
NO:			123
	77.000.40	WAY A GO OFF A COLD ADD CORED !	132
596	PR00049	WILM'S TUMOUR PROTEIN SIGNATURE	PR00049D 0.00 3.136e-09 31-46
597	DM00547	1 kw CHROMO BROMODOMAIN	DM00547C 17.30 1.667e-19 207-
		SHADOW GLOBAL.	229 DM00547E 13.94 6.200e-18 319-342 DM00547B 11.28
			1.000e-17 179-193 DM00547D
			11.60 9.250e-13 289-303
			DM00547F 23.43 6.727e-12 679-
			726 DM00547A 12.38 4.818e-11
			158-170
600	PD01066	PROTEIN ZINC FINGER ZINC- FINGER METAL-BINDING NU.	PD01066 19.43 1.882e-27 13-52
601	BL00192	Cytochrome b/b6 heme-ligand proteins.	BL00192A 11.90 6.400e-09 390-
			430
602	BL00936	Ribosomal protein L35 proteins.	BL00936B 27.27 8.615e-09 118- 157
603	BL00936	Ribosomal protein L35 proteins.	BL00936B 27.27 8.615e-09 118-
		VEVICE DISCOVERED AND A SECOND A SECON	157
606	PR00019	LEUCINE-RICH REPEAT SIGNATURE	PR00019B 11.36 7.300e-10 292- 306 PR00019A 11.19 5.667e-09
		SIGNATURE	323-337
607	PR00019	LEUCINE-RICH REPEAT	PR00019B 11.36 7.300e-10 292-
		SIGNATURE	306 PR00019A 11.19 5.667e-09
			323-337
608	PR00320	G-PROTEIN BETA WD-40 REPEAT	PR00320C 13.01 9.500e-12 168- 183 PR00320A 16.74 2.853e-10
		SIGNATURE	60-75 PR00320A 16.74 2.8336-10
			14-29 PR00320C 13.01 5.320e-10
			60-75 PR00320C 13.01 5.680e-10
	1		14-29 PR00320A 16.74 6.049e-09
			217-232 PR00320B 12.19 8.875e- 09 168-183
610	BL00750	Chaperonins TCP-1 proteins.	BL00750B 16.17 1.000e-40 70-
010	DE00730	Chapotoning for 1 protonis.	120 BL00750A 20.07 6.211e-37
			26-69 BL00750G 20.12 8.800e-31
			431-471 BL00750F 18.40 5.125e-
			30 370-411 BL00750E 24.59
			8.650e-29 295-332 BL00750H 21.44 1.000e-27 489-524
			BL00750C 25.65 5.345e-17 149-
			181 BL00750D 16.16 6.318e-14
			203-222
613	BL00766	Tetrahydrofolate	BL00766B 24.49 1.000e-40 142-
		dehydrogenase/cyclohydrolase proteins.	190 BL00766E 13.78 1.000e-40 322-359 BL00766C 25.86 5.500e-
			39 208-256 BL00766D 17.05
			4.536e-26 283-313 BL00766A
			21.48 6.063e-24 102-132
615	BL00256	Adipokinetic hormone family proteins.	BL00256 12.28 3.298e-10 746-755
616	BL00319	Amyloidogenic glycoprotein extracellular domain proteins.	BL00319C 17.12 9.053e-09 419- 453
617	BL00030	Eukaryotic RNA-binding region RNP-1 proteins.	BL00030A 14.39 4.429e-09 44-63
618	BL00030	Eukaryotic RNA-binding region RNP-1	BL00030A 14.39 4.429e-09 44-63
620	BL00325	proteins. Actin-depolymerizing proteins.	BL00325B 21.66 5.817e-16 77-
			123
622	BL00972	Ubiquitin carboxyl-terminal hydrolases	BL00972A 11.93 5.500e-19 213-

D NO. NO. Family 2 proteins. 231 BL00972D 22.55 2.742e-16 501-526 BL00972B 9.45 1.0009 11 297-307 BL00972D (2.648 3.160e-11 370-385 BL00972E 20.72 7.517e-10 526-548 3.160e-11 370-385 BL00972E 20.72 7.517e-10 526-548 3.160e-11 370-385 BL00972E 20.72 7.517e-10 526-548 PD01066 19.43 6.333e-39 6-45 PD01066 19.43 6.333e-39 6-45 PD01066 19.43 6.333e-39 6-45 PD01066 19.43 6.333e-39 6-45 PD01066 19.43 6.333e-39 6-45 PD01066 19.43 6.333e-39 6-45 PD01066 19.43 6.333e-39 6-45 PD01066 19.43 6.333e-39 6-45 PD01066 19.42 6.232-10.852 PD01066 19.42 6.232-10.852 PD01066 19.42 6.232-10.852 PD01066 19.43 6.233e-39 6-45 PD01066 19.22 8.009-12 232-246 PD01066 19.22 8.009-12 232-246 PD01066 19.22 8.009-12 232-246 PD01066 19.22 8.009-12 232-246 PD01066 19.22 8.009-12 232-246 PD01066 19.22 8.009-12 232-246 PD01066 19.22 8.009-12 232-246 PD01066 19.22 8.009-12 232-246 PD01066 19.22 8.009-12 232-246 PD01066 19.22 8.009-12 232-246 PD01066 19.22 8.009-12 232-246 PD01066 19.22 8.009-14 485-108 PD01066 19.22 8.009-14 413-409-14 63-36 PD01066 19.22 8.009-14 413-409-14 63-36 PD01066 19.22 8.009-14 413-409-14 63-36 PD01066 19.22 8.009-14 413-409-14 63-36 PD01066 19.22 8.009-14 413-409-14 63-36 PD01066 19.22 8.009-14 413-409-14 63-36 PD01066 19.22 8.009-14 413-409-14 63-36 PD01066 19.22 8.009-14 413-409-14 63-36 PD01066 19.22 8.009-14 413-409-14 63-36 PD01066 19.22 8.009-14 413-409-14 63-36 PD01066 19.22 8.009-14 413-409-14 63-36 PD01066 19.22 8.009-14 413-409-14 63-36 PD01066 19.22 8.009-14 413-409-14 63-36 PD01066 19.22 8.009-14 413-409-14 63-36 PD01066 19.22 8.009-14 413-409-14 63-36 PD01066 19.22 8.009-14 413-409-14 63-36 PD01066 19.22 8.009-14 413-409-14 63-36 PD01066 19.22 8.009-13 175-28 PD01066 19.22 8.009-13 175-28 PD01066 19.22 8.009-13 175-28 PD01066 19.22 8.009-13 175-28 PD01066 19.22 8.009-13 175-28 PD01066 19.22 8.009-13 175-28 PD01066 19.22 8.009-13 175-28 PD01066 19.22 8.009-13 175-28 PD01066 19.22 8.009-1				
family 2 proteins. 231 BL00972D 22.55 2.742e-16	ID		DESCRIPTION	RESULTS*
FINGER METAL-BINDING NU.			family 2 proteins.	501-526 BL00972B 9.45 1.000e- 11 297-307 BL00972C 16.48 3.160e-11 370-385 BL00972E
helicases proteins. S24 BL00039A 18.44 2.000e-25 198-237 BL00039C 15.63 1.844e-15 327-351 BL00039P 19.19 5.636e-14 242-268 198-237 BL00039C 15.63 1.844e-15 327-351 BL00039P 19.19 5.636e-14 242-268 PROTEIN GLYCOPROTEIN PD00306A 10.26 7.000e-12 232-246 PD00306A 10.26 7.000e-12 232-246 PROTEIN GLYCOPROTEIN PD00306A 10.26 7.000e-12 290-304 PRECURSOR RE. 304 PROUNTS STANDARD STAND	625	PD01066	I 1	
PRECURSOR RE. 246	628	BL00039	-	524 BL00039A 18.44 2.000e-25 198-237 BL00039C 15.63 1.844e- 15 327-351 BL00039B 19.19
PRECURSOR RE. 304	630	PD00306		
122 BL00785E 15,85 4,000e-16	631	PD00306	I *	
108	633	BL00785	5'-nucleotidase proteins.	122 BL00785E 15.85 4.000e-16 279-295 BL00785A 9.73 6.500e- 14 29-40 BL00785B 10.65 5.500e-13 72-86 BL00785D 9.89
DOMAIN SIGNATURE 240	636	PR00832	PAXILLIN SIGNATURE	
PR00860 VERTEBRATE METALLOTHIONEIN PR00860B 7.04 1.900e-18 85-99 PR00860C 9.61 1.474e-14 99-109 PR00860A 5.46 1.720e-14 63-76 PR00860A 5.46 1.720e-14 63-76 PD00066 13.92 4.462e-15 271-284 PD00066 13.92 2.402e-15 299-312 PD00066 13.92 2.800e-14 11-424 PD00066 13.92 2.800e-14 413-434 PD00066 13.92 2.800e-14 413-434 PD00066 13.92 2.800e-14 413-536 PD00066 13.92 8.800e-14 4495-508 PD00066 13.92 8.800e-14 4495-508 PD00066 13.92 8.800e-14 4495-508 PD00066 13.92 8.900e-13 233-536 PD00066 13.92 9.500e-13 233-536 PD00066 13.92 9.500e-13 232-536 PD00066 13.92 9.500e-13 232-536 PD00066 13.92 9.500e-13 232-536 PD00066 13.92 9.500e-13 233-536 PD00066 13.92 9.500e-13 243-256 PD00066 13.92 9.500e-13 243-256 PD00066 13.92 9.500e-13 243-256 PD00066 13.92 9.500e-13 243-256 PD00066 13.92 1.600e-09 187-200 PD00	637	PR00109	DOMAIN SIGNATURE	240
SIGNATURE PR00860C 9.61 1.474e-14 99-109 PR00860A 5.46 1.720e-14 63-76 PR07TEIN ZINC-FINGER METAL- PD00066 13.92 4.462e-15 271-284 PD00066 13.92 4.462e-15 271-284 PD00066 13.92 2.800e-14 327-340 PD00066 13.92 2.800e-14 337-394 PD00066 13.92 2.800e-14 383-396 PD00066 13.92 2.800e-14 411-424 PD00066 13.92 7.000e-14 355-368 PD00066 13.92 8.800e-14 439-452 PD00066 13.92 8.800e-14 439-452 PD00066 13.92 8.800e-13 4495-508 PD00066 13.92 7.000e-13 551-564 PD00066 13.92 7.000e-13 523-536 PD00066 13.92 9.500e-13 215-228 PD00066 13.92 9.500e-13 215-228 PD00066 13.92 9.500e-13 243-256 PD00066 13.92 9.500e-13 243-256 PD00066 13.92 8.615e-10 607-620 PD00066 13.92 8.615e-10 607-620 PD00066 13.92 8.615e-10 607-620 PD00066 13.92 8.615e-10 607-620 PD00066 13.92 8.615e-10 607-620 PD00066 13.92 8.615e-10 607-620 PD00066 13.92 8.615e-10 607-620 PD00066 13.92 8.615e-10 607-620 PD00066 13.92 8.615e-10 607-620 PD00066 13.92 8.615e-10 607-620 PD00066 13.92 8.615e-10 607-620 PD00066 13.92 8.615e-10 607-620 PD00066 13.92 8.615e-10 607-620 PD00066 13.92 8.615e-10 607-620 PD00066 13.92 8.615e-10 607-620 PD00066 13.92 8.615e-10 607-620 PD00066 13.92 8.600e-13 797-592	638	PF00635	proteins.	502
BINDI. PD00066 13.92 4.462e-15 299-312 PD00066 13.92 2.800e-14 327-340 PD00066 13.92 2.800e-14 383-396 PD00066 13.92 2.800e-14 383-396 PD00066 13.92 2.800e-14 355-368 PD00066 13.92 7.000e-13 455-368 PD00066 13.92 8.800e-14 495-508 PD00066 13.92 8.800e-14 495-508 PD00066 13.92 7.000e-13 551-564 PD00066 13.92 7.000e-13 523-536 PD00066 13.92 9.500e-13 215-228 PD00066 13.92 9.500e-13 245-228 PD00066 13.92 9.500e-13 243-256 PD00066 13.92 9.500e-13 243-256 PD00066 13.92 9.500e-13 797-592 PD00066 13.92 9.500e-13 797-592 PD00066 13.92 9.500e-13 797-592 PD00066 13.92 9.500e-13 797-592 PD00066 13.92 9.500e-13 797-592 PD00066 13.92 9.500e-13 797-592 PD00066 13.92 9.500e-13 243-256 PD00066 13.92 9.500e-1	639	PR00860	SIGNATURE	PR00860C 9.61 1.474e-14 99-109 PR00860A 5.46 1.720e-14 63-76
100 BL00961A 9.90 4.079e-26 42-66 42-66 42-66 42-66 42-66 BL00585A 28.43 1.391e-40 103-155 BL00585B 18.78 3.250e-30 193-230 193-230 193-230 BL00678 Trp-Asp (WD) repeat proteins proteins. BL00678 9.67 9.400e-10 181-192 648 PR00876 NEMATODE METALLOTHIONEIN PR00876C 6.15 9.229e-09 112-126 126 PROTEIN ZINC FINGER ZINC-FINGER METAL-BINDING NU.		·	BINDI.	PD00066 13.92 4.462e-15 299-312 PD00066 13.92 2.800e-14 327-340 PD00066 13.92 2.800e-14 383-396 PD00066 13.92 2.800e-14 411-424 PD00066 13.92 7.000e-14 355-368 PD00066 13.92 8.800e-14 439-452 PD00066 13.92 8.800e-14 495-508 PD00066 13.92 1.500e-13 551-564 PD00066 13.92 7.000e-13 467-480 PD00066 13.92 7.000e-13 523-536 PD00066 13.92 9.500e-13 215-228 PD00066 13.92 9.500e-13 243-256 PD00066 13.92 9.500e-13 579-592 PD00066 13.92 8.615e-10 607-620 PD00066 13.92 1.600e-09 187-200
155 BL00585B 18.78 3.250e-30 193-230 193-230 193-230 193-230 193-230 193-230 193-230 193-230 193-230				100 BL00961A 9.90 4.079e-26 42-66
648 PR00876 NEMATODE METALLOTHIONEIN SIGNATURE PR00876C 6.15 9.229e-09 112- 126 652 PD01066 PROTEIN ZINC FINGER ZINC- FINGER METAL-BINDING NU. PD01066 19.43 5.941e-27 29-68	643			155 BL00585B 18.78 3.250e-30 193-230
648 PR00876 NEMATODE METALLOTHIONEIN SIGNATURE PR00876C 6.15 9.229e-09 112- 126 652 PD01066 PROTEIN ZINC FINGER ZINC- FINGER METAL-BINDING NU. PD01066 19.43 5.941e-27 29-68	647	BL00678	Trp-Asp (WD) repeat proteins proteins.	BL00678 9.67 9.400e-10 181-192
FINGER METAL-BINDING NU.		L	NEMATODE METALLOTHIONEIN	126
653 BL00047 Histone H4 proteins. BL00047A 13.53 1.000e-40 2-41	652	PD01066		
	653	BL00047	Histone H4 proteins.	BL00047A 13.53 1.000e-40 2-41

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SEQ	ACCESSION	DESCRIPTION	RESULTS*
ID	NO.		10000220
NO:		}	
			BL00047B 6.51 1.429e-40 41-74
			BL00047C 12.18 1.310e-38 74-
			104
654	PD01066	PROTEIN ZINC FINGER ZINC-	PD01066 19.43 4.109e-25 30-69
	L	FINGER METAL-BINDING NU.	
655	BL01115	GTP-binding nuclear protein ran proteins.	BL01115A 10.22 3.483e-17 19-63
657	BL00518	Zinc finger, C3HC4 type (RING finger), proteins.	BL00518 12.23 8.286e-10 31-40
658	BL00125	Serine/threonine specific protein	BL00125B 21.48 1.000e-40 89-
		phosphatases proteins.	135 BL00125C 19.97 1.000e-40
			153-200 BL00125D 33.11 1.000e-
			40 213-268 BL00125A 14.83
- (20	7700000	DROWERS AND EDICED METAL	8.941e-38 47-84 PD00066 13.92 8.200e-16 492-505
659	PD00066	PROTEIN ZINC-FINGER METAL-	PD00066 13.92 8.200e-16 492-303 PD00066 13.92 9.308e-15.380-393
		BINDI.	PD00066 13.92 9.306e-13.380-393
			PD00066 13.92 7.000e-13 240-253
			PD00066 13.92 7.500e-13 268-281
			PD00066 13.92 7.500e-13 408-421
			PD00066 13.92 2.174e-11 464-477
		·	PD00066 13.92 1.000e-10 436-449
660	PD01066	PROTEIN ZINC FINGER ZINC-	PD01066 19.43 2.189e-26 29-68
		FINGER METAL-BINDING NU.	
661	BL00795	Involucrin proteins.	BL00795C 17.06 7.882e-15 193-
			238 BL00795C 17.06 3.797e-13
			187-232 BL00795C 17.06 5.014e-
			13 188-233 BL00795C 17.06
			4.506e-12 196-241 BL00795C
1			17.06 7.896e-12 191-236
		X X	BL00795C 17.06 1.667e-11 185-
			230 BL00795C 17.06 2.000e-11
			198-243 BL00795C 17.06 3.778e-
			11 171-216 BL00795C 17.06
			6.111e-11 197-242 BL00795C
			17.06 6.444e-11 194-239 BL00795C 17.06 8.000e-11 189-
			234 BL00795C 17.06 8.000e-11 189-
•			192-237 BL00795C 17.06 1.733e-
			10 195-240 BL00795C 17.06
			2.779e-10 184-229 BL00795C
		•	17.06 4.035e-10 199-244
			BL00795C 17.06 5.081e-10 186-
		.}	231 BL00795C 17.06 6.965e-10
		1	190-235 BL00795C 17.06 2.700e-
			09 200-245 BL00795C 17.06
		1	5.800e-09 175-220 BL00795C
			17.06 6.500e-09 182-227
			BL00795C 17.06 6.600e-09 201-
			246 BL00795C 17.06 6.600e-09
]		j	202-247 BL00795C 17.06 6.600e-
			09 208-253
662	BL00469	Nucleoside diphosphate kinases proteins.	BL00469 22.22 1.000e-40 149-204
663	BL01160	Kinesin light chain repeat proteins.	BL01160B 19.54 9.411e-11 331-
			385
664	BL00601	Tryptophan pentad repeat proteins (IRF	BL00601A 20.29 5.500e-23 7-46
		family) proteins.	BL00601B 20.92 3.631e-13 69-98
665	BL00082	Extradiol ring-cleavage dioxygenases	BL00082A 19.07 8.615e-12 49-72
	70.00	proteins.	D) (01527D 01 (2 4 072 - 27 074
666	DM01537	kw SKI2W SKI2 NUCLEOLAR	DM01537B 21.63 4.073e-37 834-

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SEQ ID NO:	ACCESSION NO.	DESCRIPTION	RESULTS*
		HELICASE.	881 DM01537B 21.63 9.750e-21 1669-1716 DM01537A 15.14 8.650e-18 698-718 DM01537A 15.14 6.766e-12 1537-1557
667	DM01537	kw SKI2W SKI2 NUCLEOLAR HELICASE.	DM01537B 21.63 7.923e-38 820- 867 DM01537B 21.63 9.750e-21 1655-1702 DM01537A 15.14 8.650e-18 684-704 DM01537A 15.14 6.766e-12 1523-1543
669	BL00107	Protein kinases ATP-binding region proteins.	BL00107A 18.39 6.786e-24 849- 880 BL00107B 13.31 6.727e-13 916-932
670	BL00299	Ubiquitin domain proteins.	BL00299 28.84 9.735e-27 37-89
671	BL00027	'Homeobox' domain proteins.	BL00027 26.43 6.571e-12 432-475
676	PR00861	ALPHA-LYTIC ENDOPEPTIDASE SERINE PROTEASE (S2A) SIGNATURE	PR00861E 9.88 2.385e-09 206- 221
678	BL00225	Crystallins beta and gamma 'Greek key' motif proteins.	BL00225B 18.06 7.517e-24 1805- 1840 BL00225B 18.06 8.297e-20 1987-2022 BL00225B 18.06 2.575e-19 1896-1931 BL00225B 18.06 8.200e-19 175-210 BL00225B 18.06 8.200e-19 1698- 1733 BL00225B 18.06 4.808e-14 73-108 BL00225B 18.06 4.808e- 14 1596-1631 BL00225B 18.06 5.500e-14 2077-2112 BL00225A 13.82 5.829e-12 2043-2064 BL00225A 13.82 3.127e-09 1759- 1780
679	PR00320	G-PROTEIN BETA WD-40 REPEAT SIGNATURE	PR00320C 13.01 4.240e-10 169- 184 PR00320A 16.74 6.294e-10 169-184
680	BL00243	Integrins beta chain cysteine-rich domain proteins.	BL00243I 31.77 1.143e-11 172- 215
681	PR00852	XERODERMA PIGMENTOSUM GROUP D PROTEIN SIGNATURE	PR00852H 5.90 1.000e-29 612-635 PR00852E 8.14 3.769e-27 348-371 PR00852D 11.38 8.875e-27 309-331 PR00852B 11.08 2.800e-25 249-269 PR00852I 17.26 3.500e-25 683-704 PR00852F 11.85 5.909e-24 379-398 PR00852G 16.19 4.462e-23 468-486 PR00852C 8.81 9.143e-23 284-303
682	BL50058	G-protein gamma subunit profile.	BL50058 27.23 1.375e-35 15-63
685	BL00972	Ubiquitin carboxyl-terminal hydrolases family 2 proteins.	BL00972A 11.93 7.500e-20 40-58 BL00972D 22.55 3.903e-16 300- 325 BL00972B 9.45 1.000e-13 120-130 BL00972E 20.72 5.500e- 11 325-347
687	BL00237	G-protein coupled receptors proteins.	BL00237A 27.68 4.273e-14 98- 138
688	BL00388	Proteasome A-type subunits proteins.	BL00388A 23.14 1.000e-40 8-54 BL00388B 31.38 3.864e-33 66- 108 BL00388D 20.71 1.000e-21 153-184 BL00388C 18.79 8.147e-

PROTEIN STEROL CARRIER LIPID-

PD02796

689

16 126-148 PD02796B 20.92 1.105e-15 347-

CEO	ACCESSION	DESCRIPTION	RESULTS*
SEQ	NO.		ACESCE 15
NO:	110.		
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691	PD01572	PHOTOSYSTEM II REACTION	PD01572 8.77 4.083e-09 1-31
"-		CENTRE T PROTEIN PHOTOS.	
692	BL00028	Zinc finger, C2H2 type, domain proteins.	BL00028 16.07 7.600e-10 488-505
694	BL01013	Oxysterol-binding protein family	BL01013A 25.14 9.357e-33 527-
		proteins.	563 BL01013D 26.81 8.235e-23
			814-858 BL01013C 9.97 6.211e-
			14 615-625 BL01013B 11.33
			3.605e-13 592-603
695	PD00289	PROTEIN SH3 DOMAIN REPEAT	PD00289 9.97 3.571e-13 164-178
		PRESYNA.	PD00289 9.97 8.650e-11 2147- 2161 PD00289 9.97 2.552e-09 23-
			37
698	PR00161	NICKEL-DEPENDENT	PR00161C 9.51 4.930e-09 282-
098	PROOTOT	HYDROGENASE/B-TYPE	302
		CYTOCHROME SIGNATURE	302
700	PR00749	LYSOZYME G SIGNATURE	PR00749F 13.63 8.636e-13 139-
'**	1100715	2.50212 0 0.01	156 PR00749H 8.22 3.681e-12
			173-194 PR00749B 16.54 1.419e-
			11 48-70 PR00749C 7.26 3.060e-
			11 72-91 PR00749A 10.33
·			4.815e-10 24-45
703	PR00704	CALPAIN CYSTEINE PROTEASE (C2)	PR00704I 9.52 1.000e-29 476-505
		FAMILY SIGNATURE	PR00704D 11.05 2.500e-27 132-
1			158 PR00704E 12.55 5.500e-27
1			162-186 PR00704F 13.61 1.000e- 22 187-215 PR00704G 13.87
			1.237e-21 317-339 PR00704H
		·	13.38 8.138e-21 367-385
,			PR00704A 14.68 2.125e-19 27-51
		4	PR00704C 11.88 1.257e-17 96-
			113 PR00704B 17.94 1.833e-15
1			72-95
705	PR00859	PROKARYOTE METALLOTHIONEIN	PR00859C 7.06 2.776e-09 94-111
		SIGNATURE	
706	BL00226	Intermediate filaments proteins.	BL00226D 19.10 9.581e-26 369-
			416 BL00226B 23.86 3.250e-24
			203-251 BL00226C 13.23 8.269e-
			21 268-299 BL00226A 12.77 8.200e-14 103-118
707	PR00021	SMALL PROLINE-RICH PROTEIN	PR00021A 4.31 2.440e-10 2-15
707	PR00021	SIGNATURE	FR00021A 4.51 2.4406-10 2-15
708	BL00361	Ribosomal protein S10 proteins.	BL00361B 18.34 5.101e-10 82-
'00	DECOSOT	recording process of process.	105
709	PR00021	SMALL PROLINE-RICH PROTEIN	PR00021A 4.31 2.200e-10 2-15
}	11000	SIGNATURE	
710	BL00514	Fibrinogen beta and gamma chains C-	BL00514C 17.41 8.412e-27 160-
i		terminal domain proteins.	197 BL00514E 14.28 8.909e-16
		_	219-236 BL00514H 14.95 1.551e-
<u> </u>			15 317-342 BL00514G 15.98
			7.750e-15 284-314 BL00514D
			15.35 4.789e-10 201-214
711	PD00930	PROTEIN GTPASE DOMAIN	PD00930B 33.72 8.714e-12 49-90
	77.00.400	ACTIVATION.	77.00400004.50.6000 45.150
714	BL00400	LBP / BPI / CETP family proteins.	BL00400C 24.53 6.029e-17 158-
			202 BL00400D 23.26 2.080e-14
			222-259 BL00400A 21.59 1.600e- 10 27-59
715	BL01154	RNA polymerases L / 13 to 16 Kd	BL01154B 24.55 5.500e-36 40-76
(17)	DL01134	INTA POLYMETASES L / 13 to 10 Ku	DE0113-D 24.33 3.3006-30 40-70

SEQ ID NO:	ACCESSION NO.	DESCRIPTION	RESULTS*
		subunits proteins.	BL01154A 18.70 3.000e-22 19-40
716	PD01066	PROTEIN ZINC FINGER ZINC- FINGER METAL-BINDING NU.	PD01066 19.43 9.786e-32 10-49
717	BL00215	Mitochondrial energy transfer proteins.	BL00215A 15.82 9.206e-14 77- 102 BL00215A 15.82 8.412e-10 175-200
719	BL00309	Vertebrate galactoside-binding lectin proteins.	BL00309C 18.65 2.241e-09 62-87
726	BL00687	Aldehyde dehydrogenases glutamic acid proteins.	BL00687E 25.37 7.136e-33 266- 316 BL00687D 26.00 5.333e-28 151-198 BL00687B 17.54 3.647e- 26 39-81 BL00687C 24.13 6.087e-22 96-133 BL00687F 9.55 2.500e-11 352-363
727	DM01354	kw TRANSCRIPTASE REVERSE II ORF2.	DM01354N 13.17 1.000e-40 129- 174 DM01354O 8.73 6.605e-15 180-226
734	PD00301	PROTEIN REPEAT MUSCLE CALCIUM-BI.	PD00301A 10.24 6.400e-09 101- 112
735	BL01024	Protein phosphatase 2A regulatory subunit PR55 proteins.	BL01024A 10.26 1.000e-40 22-69 BL01024B 8.91 1.000e-40 86-127 BL01024C 7.80 1.000e-40 146- 185 BL01024D 13.22 1.000e-40 185-222 BL01024E 11.96 1.000e- 40 222-266 BL01024F 9.42 1.000e-40 266-317 BL01024G 11.09 1.000e-40 317-349 BL01024H 13.88 1.000e-40 389- 442
736	PF00913	Trypanosome variant surface glycoprotein.	PF00913D 11.90 7.130e-10 24-51
737	PR00700	PROTEIN TYROSINE PHOSPHATASE SIGNATURE	PR00700D 12.47 2.200e-09 82- 101
740	PR00320	G-PROTEIN BETA WD-40 REPEAT SIGNATURE	PR00320C 13.01 1.600e-09 68-83 PR00320A 16.74 7.366e-09 68-83
743	PR00871	DNA NUCLEOTIDYLEXOTRANSFERASE (TDT) SIGNATURE	PR00871G 14.48 8.000e-09 178- 201
745	BL00518	Zinc finger, C3HC4 type (RING finger), proteins.	BL00518 12.23 2.286e-10 33-42
749	BL00215	Mitochondrial energy transfer proteins.	BL00215A 15.82 5.200e-15 221- 246 BL00215A 15.82 7.618e-14 20-45 BL00215A 15.82 8.851e-11 123-148 BL00215B 10.44 9.526e- 11 69-82 BL00215B 10.44 7.300e-09 272-285 BL00215B 10.44 8.500e-09 165-178
751	BL50002	Src homology 3 (SH3) domain proteins profile.	BL50002A 14.19 1.000e-14 370- 389 BL50002B 15.18 2.200e-10 408-422
752	BL00353	HMG1/2 proteins.	BL00353B 11.47 3.089e-12 390- 440
753	PF00622	Domain in SPIa and the RYanodine Receptor.	PF00622B 21.00 4.214e-14 47-69
754	BL00211	ABC transporters family proteins.	BL00211A 12.23 8.941e-10 66-78
755	PR00926	MITOCHONDRIAL CARRIER PROTEIN SIGNATURE	PR00926F 17.75 7.750e-19 392- 415 PR00926C 16.07 5.935e-17 253-274 PR00926D 10.53 2.059e- 15 301-320 PR00926E 11.70

CEO	ACCESSION	DESCRIPTION	RESULTS*
SEQ	NO.	DESCRIE I TON	1000010
NO:	110.		
			4.971e-15 344-363 PR00926B
			16.07 9.526e-13 210-225
			PR00926A 10.41 1.514e-12 197-
	5101107	California FOR No. 1	211 BL01187A 9.98 2.125e-12 324-
756	BL01187	Calcium-binding EGF-like domain proteins pattern proteins.	336 BL01187A 9.98 4.789e-11
]		proteins pattern proteins.	377-389 BL01187B 12.04 3.057e-
			10 439-455
757	PF00651	BTB (also known as BR-C/Ttk) domain	PF00651 15.00 4.429e-10 43-56
		proteins.	
758	PR00055	HIV TAT DOMAIN SIGNATURE	PR00055A 8.13 8.855e-09 144-
		The second control of the second seco	156
759	PD00066	PROTEIN ZINC-FINGER METAL-	PD00066 13.92 5.304e-11 110-123
760	PR00448	BINDI. NSF ATTACHMENT PROTEIN	PR00448D 12.42 3.455e-27 162-
/60	PK00448	SIGNATURE°	186 PR00448A 10.74 1.273e-22
		BIGINITORE	37-57 PR00448B 16.01 9.379e-21
			100-118 PR00448C 11.46 1.000e-
]			20 129-147
765	BL01042	Homoserine dehydrogenase proteins.	BL01042A 13.29 5.909e-11 74-95
766	PR00625	DNAJ PROTEIN FAMILY	PR00625A 12.84 2.154e-18 26-46
760	DI 007/2	SIGNATURE WHEP-TRS domain proteins.	PR00625B 13.48 9.000e-16 57-78 BL00762A 23.43 8.500e-28 112-
768	BL00762	WHEP-1RS domain proteins.	149 BL00762B 16.14 3.793e-12
			64-78 BL00762A 23.43 6.625e-12
			6-43 BL00762C 15.58 4.176e-09
			459-472 BL00762D 11.15 9.667e-
			09 210-220
769	PR00709	AVIDIN SIGNATURE	PR00709A 4.60 1.934e-09 1-20
770	PR00320	G-PROTEIN BETA WD-40 REPEAT	PR00320C 13.01 1.720e-10 262- 277 PR00320A 16.74 2.853e-10
		SIGNATURE	262-277 PR00320C 13.01 4.300e-
			09 96-111 PR00320B 12.19
		•	5.500e-09 262-277 PR00320A
			16.74 6.268e-09 55-70
771	PR00019	LEUCINE-RICH REPEAT	PR00019B 11.36 8.714e-12 87-
		SIGNATURE	101 PR00019A 11.19 1.000e-10
	770000	A DOLIDODO CERTIFE PRECIPION	90-104 PD02807C 8.91 6.308e-10 110-
772	PD02807	APOLIPOPROTEIN E PRECURSOR APO-E GLYCOPROTEIN PLAS.	159
773	PD02807	APOLIPOPROTEIN E PRECURSOR	PD02807C 8.91 6.308e-10 155-
['''	1 102001	APO-E GLYCOPROTEIN PLAS.	204
774	DM00547	1 kw CHROMO BROMODOMAIN	DM00547F 23.43 3.942e-28 943-
		SHADOW GLOBAL.	990 DM00547E 13.94 9.750e-21
[652-675 DM00547B 11.28
			1.818e-18 518-532 DM00547C
[17.30 3.531e-17 546-568 DM00547A 12.38 1.273e-11 497-
			509 DM00547D 11.60 9.200e-11
			622-636
776	PR00779	INOSITOL 1,4,5-TRISPHOSPHATE-	PR00779F 14.51 5.147e-09 769-
		BINDING PROTEIN RECEPTOR	792
		SIGNATURE	
777	PR00779	INOSITOL 1,4,5-TRISPHOSPHATE-	PR00779F 14.51 5.147e-09 742-
	•	BINDING PROTEIN RECEPTOR	765
	DE 00.550	SIGNATURE	DD00770F 14 51 5 147- 00 742
778	PR00779	INOSITOL 1,4,5-TRISPHOSPHATE- BINDING PROTEIN RECEPTOR	PR00779F 14.51 5.147e-09 742- 765
		SIGNATURE	103
		PIGNATOKE	L

PF00084B 9.45 6.400e-09 656-668				The state of the s
NO:	-		DESCRIPTION	RESULTS*
779 BL01282 BIR repeat proteins. BL01282B 30.49 2.543e-09 6-45		NO.	,	
PR00205		BI 01282	BIR repeat proteins	BI 01282B 30 49 2 543e-09 6-45
Section				
230-248 PR00205B 11.39 8.527e- 10 551-559 PR00205B 11.39 8.527e- 10 551-559 PR00205B 11.39 8.527e- 10 551-559 PR00205B 11.39 8.527e- 10 551-559 PR00205B 11.39 8.527e- 10 551-559 PR00205B 11.39 8.527e- 11 59-228 BL.00625B 17.69 1.885e- 16 140-174 BL.00625B 17.69 1.885e- 16 140-174 BL.00625B 17.69 1.885e- 16 140-174 BL.00625B 17.69 1.885e- 16 140-174 BL.00625B 17.69 1.885e- 16 140-174 BL.00625B 17.69 1.885e- 16 140-174 BL.00625B 17.69 1.885e- 16 140-174 BL.00625B 17.69 1.885e- 16 140-174 BL.00625B 17.69 1.885e- 16 140-174 BL.00625B 17.69 1.885e- 16 140-174 BL.00625B 17.69 1.885e- 17 57	/01	1100203	O'B'IB'G' O'G' WI'O'G	
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Regulator of chromosome condensation (RCC1) proteins. BL00625B 17.69 2.1676-19 193-227 BL00625A 16.21 5.500e-17 199-228 BL00625B 17.69 1.885e-16 140-174 BL00625B 17.69 1.885e-16 140-174 BL00625B 17.69 1.885e-16 140-174 BL00625B 17.69 1.885e-16 140-174 BL00625B 17.69 1.885e-16 140-174 BL00625B 17.69 1.885e-16 140-174 BL00625B 17.69 1.885e-16 140-174 BL00625B 17.69 1.885e-16 140-174 BL00625B 17.69 1.885e-16 140-174 BL00625B 17.69 1.885e-16 1507e-14 146-175				10 551-569 PR00205B 11.39
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A DOMAIN SIGNATURE	767	DL00820 .	WARCES failing proteins.	
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A37 BL00415N 4.29 2.117e-09 103-147 BL00415N 4.29 3.628e-			proteins.	209
103-147 BL00415N 4.29 3.628e- 09 97-141 BL00415N 4.29 5.664e-09 387-431 FOUR PROTEIN ZINC FINGER ZINC-FINGER METAL-BINDING NU. PD01066	791	BL00415	Synapsins proteins.	7
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380 PF00731B 19.47 7.429e-28 299-336 PF00731A 19.32 6.333e-24 268-297	·			
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BL00170 Cyclophilin-type peptidyl-prolyl cis-trans isomerase signatur. 337 337 337 337 337 BL00678 Trp-Asp (WD) repeat proteins proteins. BL00678 9.67 3.400e-10 378-389 BL00678 9.67 5.800e-10 418-429 BL00678 9.67 8.800e-10 418-429 BL00678 9.67 8.800e-10 295-306 B06 PD01719 PRECURSOR GLYCOPROTEIN PD01719A 12.89 7.571e-14 290- SIGNAL RE. 318 PR00320 G-PROTEIN BETA WD-40 REPEAT PR00320B 12.19 9.100e-09 451- 466 BL00107 Protein kinases ATP-binding region BL00107A 18.39 4.462e-12 564- proteins. Protein kinases ATP-binding region PR00453 VON WILLEBRAND FACTOR TYPE PR00453A 12.79 1.310e-14 36-54 A DOMAIN SIGNATURE PR00453B 14.65 8.568e-10 75-90 PROTEIN ZINC FINGER ZINC- FINGER METAL-BINDING NU. PD01066 19.43 2.047e-31 16-55 FINGER METAL-BINDING NU. PR00193D 14.36 5.154e-36 125- 154 PR00193E 19.47 3.919e-18 179-208 PR00193E 19.47 3.919e-18 179-208 PR00193E 19.47 3.919e-18 PR00193E 19				· ·
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BL00678 Trp-Asp (WD) repeat proteins proteins. BL00678 9.67 3.400e-10 378-389	30-r	2200110		
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Recursor Glycoprotein PD01719A 12.89 7.571e-14 290-318			•	1
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PR00320 G-PROTEIN BETA WD-40 REPEAT PR00320B 12.19 9.100e-09 451- 466 SIGNATURE 466 BL00107 Protein kinases ATP-binding region proteins. BL00107A 18.39 4.462e-12 564- 595 SIGNATURE PR00453A 12.79 1.310e-14 36-54 PR00453B 14.65 8.568e-10 75-90 PROTEIN ZINC FINGER ZINC- FINGER METAL-BINDING NU. PD01066 19.43 2.047e-31 16-55 PD01066 PROTEIN ZINC FINGER ZINC- FINGER METAL-BINDING NU. PR00193D 14.36 5.154e-36 125- SIGNATURE PR00193E 19.47 3.919e-18 179-208 PR00193E 19.47 3.919e-18 PR00193E 19.47 3.919E-18 PR00193E 19.47 3.919E-18 PR00193E 19.47 3.919E-18 PR00193E 19.47 3.	806	PD01719		
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A DOMAIN SIGNATURE			proteins.	595
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FINGER METAL-BINDING NU. 817 PR00193 MYOSIN HEAVY CHAIN PR00193D 14.36 5.154e-36 125- SIGNATURE 154 PR00193E 19.47 3.919e-18 179-208	815	PDOTORE		PD01066 19 43 2 0476 31 16 55
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179-208	817	PR00193	MYOSIN HEAVY CHAIN	
			SIGNATURE	
616 PROUSSU ENDUPERTIDASE LA (LON) SERINE PROUSSUA 8.41 9.5716-11 115-	910	DDOOGG	PMDODERATE A GOVERNOR	1
	618	PK00830	ENDOPERTIDASE LA (LON) SERINE	PKUU83UA 8.41 9.3/16-11 115-

SEQ	ACCESSION	DESCRIPTION	RESULTS*
ID	NO.		
NO:			
		PROTEASE (S16) SIGNATURE	135
819	BL00126	3'5'-cyclic nucleotide phosphodiesterases	BL00126C 22.07 7.857e-24 528-
		proteins.	569 BL00126E 35.22 3.714e-15
			669-724 BL00126D 25.50 1.173e-
1			14 584-623 BL00126B 15.20
			1.000e-12 502-514 BL00126A
			27.56 3.361e-09 461-498
820	PR00511	TEKTIN SIGNATURE	PR00511B 12.25 8.826e-22 174-
]			195 PR00511A 13.59 7.723e-11
	DI 00541		155-172
821	BL00741	Guanine-nucleotide dissociation	BL00741B 14.27 2.800e-15 13-36
	77700700	stimulators CDC24 family sign.	DD00500114 C0 4 005 - 00 021
822	PF00780	Domain found in NIK1-like kinases,	PF00780I 14.69 4.825e-09 231-
	DV 00000	mouse citron and yeast ROM.	261
827	BL00030	Eukaryotic RNA-binding region RNP-1	BL00030A 14.39 5.235e-11 144-
	77.0000	proteins.	163
828	BL00326	Tropomyosins proteins.	BL00326D 8.76 9.357e-11 545- 586
- 000	77700110	TRANSCRIPTION PROTEIN DNA-	PD02448A 9.37 1.000e-40 46-85
829	PD02448	BINDIN.	PD02448B 10.17 1.000e-40 46-85
}		BINDIN.	133 PD02448C 13.62 1.000e-40
1.			152-189 PD02448E 11.33 9.000e-
			30 235-261 PD02448F 14.22
			9.654e-25 279-303 PD02448D
			11.48 3.659e-18 197-211
			PD02448G 10.73 7.857e-16 305-
			318
830	BL00720	Guanine-nucleotide dissociation	BL00720B 16.57 4.500e-23 483-
030	DD00120	stimulators CDC25 family sign.	507
831	BL00107	Protein kinases ATP-binding region	BL00107A 18.39 6.625e-21 143-
	BECOTO	proteins.	174 BL00107B 13.31 4.214e-10
		proteins	213-229
832	BL00215	Mitochondrial energy transfer proteins.	BL00215A 15.82 5.787e-11 32-57
833	PR00497	NEUTROPHIL CYTOSOL FACTOR	PR00497A 6.92 4.375e-09 41-59
		P40 SIGNATURE	
834	BL00229	Tau and MAP proteins tubulin-binding	BL00229A 23.57 9.565e-10 99-
		domain proteins.	138
835	BL00421	Transmembrane 4 family proteins.	BL00421E 20.97 2.216e-09 1053-
			1083
836	BL00795	Involucrin proteins.	BL00795B 12.41 7.931e-09 405-
			445
837	PR00020	MAM DOMAIN SIGNATURE	PR00020A 18.17 1.000e-17 34-53
			PR00020B 15.52 5.846e-16 68-85
		•	PR00020D 12.70 2.543e-15 147-
			162 PR00020C 13.66 3.483e-13
			95-107 PR00020E 8.64 6.586e-13
			165-179
838	BL50017	Death domain proteins profile.	BL50017B 17.60 6.897e-13 1499-
			1515
839	PF00850	Histone deacetylase family.	PF00850C 14.55 9.542e-09 1352-
			1369
840	PF00023	Ank repeat proteins.	PF00023A 16.03 4.500e-12 44-60
			PF00023B 14.20 7.923e-11 73-83
			PF00023B 14.20 9.000e-10 139-
			149 PF00023B 14.20 5.500e-09
			40-50
842	BL01194	Ribosomal protein L15e proteins.	BL01194B 13.66 1.000e-40 37-85
[[BL01194C 12.35 9.250e-40 103-
		<u> </u>	138 BL01194A 18.70 7.632e-38

SEQ	ACCESSION	DESCRIPTION	RESULTS*
ID NO:	NO.		
			2-37 BL01194D 19.02 2.658e-36 139-178
843	BL00610	Sodium:neurotransmitter symporter family proteins.	BL00610A 17.73 1.000e-40 40-90 BL00610B 23.65 1.000e-40 104- 154 BL00610C 12.94 1.000e-40 206-258 BL00610E 20.34 1.000e- 40 355-398 BL00610F 29.02 1.000e-40 454-509 BL00610D 20.97 6.063e-35 272-325 BL00610G 12.89 8.588e-13 514- 537
845	BL00143	Insulinase family, zinc-binding region proteins.	BL00143A 20.91 4.300e-20 94- 121 BL00143C 14.16 5.500e-13 245-258 BL00143B 14.41 9.053e- 10 141-156
846	PR00543	OESTROGEN RECEPTOR SIGNATURE	PR00543D 10.87 1.355e-09 898- 914
847	PR00543	OESTROGEN RECEPTOR SIGNATURE	PR00543D 10.87 1.355e-09 898- 914
848	BL00824	Elongation factor 1 beta/beta'/delta chain proteins.	BL00824C 14.58 1.000e-40 129- 167 BL00824D 14.04 6.192e-39 167-202 BL00824B 9.21 2.080e- 21 96-116 BL00824E 12.49 3.333e-19 210-226 BL00824A 13.78 8.650e-14 19-34
849	PD01066	PROTEIN ZINC FINGER ZINC- FINGER METAL-BINDING NU.	PD01066 19.43 1.000e-40 12-51
850	PD01066	PROTEIN ZINC FINGER ZINC- FINGER METAL-BINDING NU.	PD01066 19.43 7.316e-24 10-49
852	BL01272	Glucokinase regulatory protein family proteins.	BL01272B 19.61 6.870e-30 136- 171 BL01272C 11.68 3.314e-25 249-274 BL01272A 6.49 1.231e- 18 99-117
853	PD00930	PROTEIN GTPASE DOMAIN ACTIVATION.	PD00930B 33.72 9.341e-20 65- 106
854	PD00289	PROTEIN SH3 DOMAIN REPEAT PRESYNA.	PD00289 9.97 6.850e-11 140-154
858	PR00450	RECOVERIN FAMILY SIGNATURE	PR00450C 12.22 3.250e-25 68-90 PR00450B 11.76 8.125e-23 22-42 PR00450D 16.58 8.920e-22 92- 112 PR00450E 12.14 1.581e-19 114-133 PR00450G 15.33 5.500e- 19 166-187 PR00450F 12.30 4.375e-15 140-156 PR00450A 13.58 1.857e-14 8-23
860	BL00027	'Homeobox' domain proteins.	BL00027 26.43 7.188e-27 74-117
866	BL00477	Alpha-2-macroglobulin family thiolester region proteins.	BL00477L 23.51 7.480e-20 54-87
867	BL01078	Molybdenum cofactor biosynthesis proteins.	BL01078B 14.20 1.621e-20 408- 429 BL01078A 10.16 2.000e-13 366-379 BL01078D 5.99 3.455e- 11 566-576 BL01078C 10.52 3.793e-11 501-513
868	BL01177	Anaphylatoxin domain proteins.	BL01177E 20.64 5.800e-24 462- 489 BL01177C 17.39 5.333e-19 416-435 BL01177B 13.61 7.840e- 16 122-138 BL01177D 17.50 1.900e-15 441-459
869	BL01177	Anaphylatoxin domain proteins.	BL01177E 20.64 5.800e-24 415-

SEQ ACCESSION NO. NO: BL50007 Phosphatidylinositol-spec phospholipase X-box dor prof. 872 BL00972 Ubiquitin carboxyl-termi family 2 proteins.	442 BL01177C 17.39 5.333e-19 369-388 BL01177B 13.61 7.840e- 16 122-138 BL01177D 17.50 1.900e-15 394-412 cific main proteins BL50007A 19.61 1.000e-40 322- 368 BL50007D 19.54 1.000e-40 589-631 BL50007B 20.90 6.700e- 36 383-421 BL50007E 25.63 9.053e-33 748-785 BL50007C 8.97 5.200e-19 452-469 inal hydrolases BL00972D 22.55 3.250e-17 90- 115 URE PR00452B 11.65 4.250e-09 370- 386
871 BL50007 Phosphatidylinositol-spec phospholipase X-box dor prof. 872 BL00972 Ubiquitin carboxyl-termi family 2 proteins.	369-388 BL01177B 13.61 7.840e- 16 122-138 BL01177D 17.50 1.900e-15 394-412 cific BL50007A 19.61 1.000e-40 322- 368 BL50007D 19.54 1.000e-40 589-631 BL50007B 20.90 6.700e- 36 383-421 BL50007E 25.63 9.053e-33 748-785 BL50007C 8.97 5.200e-19 452-469 inal hydrolases BL00972D 22.55 3.250e-17 90- 115 URE PR00452B 11.65 4.250e-09 370- 386
phospholipase X-box dor prof. 872 BL00972 Ubiquitin carboxyl-termi family 2 proteins.	main proteins 368 BL50007D 19.54 1.000e-40 589-631 BL50007B 20.90 6.700e- 36 383-421 BL50007E 25.63 9.053e-33 748-785 BL50007C 8.97 5.200e-19 452-469 inal hydrolases BL00972D 22.55 3.250e-17 90- 115 URE PR00452B 11.65 4.250e-09 370- 386
family 2 proteins.	115 URE PR00452B 11.65 4.250e-09 370- 386
	386
874 PR00452 SH3 DOMAIN SIGNAT	
877 BL00741 Guanine-nucleotide disso stimulators CDC24 famil	ly sign. 1366
878 DM00215 PROLINE-RICH PROTE	EIN 3. DM00215 19.43 2.525e-09 52-85
881 PD02807 APOLIPOPROTEIN E P	PRECURSOR PD02807E 10.90 4,702e-09 358-
APO-E GLYCOPROTEI	
882 PD01066 PROTEIN ZINC FINGE FINGER METAL-BIND	R ZINC- PD01066 19.43 7.188e-37 8-47
885 PF00023 Ank repeat proteins.	PF00023A 16.03 8.071e-09 10-26
886 PR00372 BIOPTERIN-DEPENDE AROMATIC AMINO A HYDROXYLASE SIGN	CID 248 PR00372A 13.39 7.000e-24
887 BL00301 GTP-binding elongation	
888 BL00518 Zinc finger, C3HC4 type proteins.	
889 PD01066 PROTEIN ZINC FINGE FINGER METAL-BIND	•
890 DM00179 w KINASE ALPHA ADI	HESION T- DM00179 13.97 7.652e-09 113- 123
892 BL01022 PTR2 family proton/oligo symporters proteins.	118 BL01022E 23.51 1.173e-12 472-508 BL01022A 11.58 9.135e- 12 42-61 BL01022D 9.42 3.455e- 11 199-212
893 PD02407 3-BISPHOSPHOGLYCE INDEPENDENT PHOSE	PHOGLYCER. 383
894 PD02407 3-BISPHOSPHOGLYCE INDEPENDENT PHOSE	PHOGLYCER. 383
895 PR00237 RHODOPSIN-LIKE GPO SUPERFAMILY SIGNA	TURE 138 PR00237F 13.57 1.360e-13 312-337 PR00237G 19.63 9.069e- 13 353-380 PR00237E 13.03 7.120e-12 243-267 PR00237D 8.94 4.150e-11 194-216 PR00237A 11.48 4.375e-11 83- 108
896 BL00129 Glycosyl hydrolases fami	ily 31 proteins. BL00129D 16.76 8.258e-26 634-678 BL00129A 26.21 1.720e-25 384-430 BL00129E 22.60 4.857e-

SEQ ID NO:	ACCESSION NO.	DESCRIPTION	RESULTS*
No.			23 698-734 BL00129C 15.12 1.750e-22 596-624 BL00129B 19.19 5.891e-18 495-522 BL00129F 26.19 7.545e-15 814- 852
897	BL00598	Chromo domain proteins.	BL00598 14.45 1.220e-13 9-31
898	BL00518	Zinc finger, C3HC4 type (RING finger),	BL00518 12.23 6.000e-09 396-405
- 000	77201101	proteins.	PD01101B 21.53 1.000e-40 274-
899	PD01101	INHIBITOR HEAVY CHAIN CHANNEL IN.	327 PD01101D 24.45 1.000e-40 457-512 PD01101A 18.25 6.268e- 23 83-117 PD01101C 12.69 1.237e-16 366-386 PD01101E 6.73 7.750e-12 566-576
900	PR00600	PROTEIN PHOSPHATASE PP2A 55KD REGULATORY SUBUNIT SIGNATURE	PR00600A 11.61 5.979e-09 31-52
901	PD01066	PROTEIN ZINC FINGER ZINC- FINGER METAL-BINDING NU.	PD01066 19.43 8.116e-31 24-63
903	BL01115	GTP-binding nuclear protein ran proteins.	BL01115A 10.22 1.509e-11 21-65
906	DM00215	PROLINE-RICH PROTEIN 3.	DM00215 19.43 2.174e-13 539- 572 DM00215 19.43 4.750e-12 549-582 DM00215 19.43 9.824e- 11 551-584 DM00215 19.43 2.929e-10 548-581 DM00215 19.43 4.054e-10 550-583 DM00215 19.43 5.339e-10 552- 585 DM00215 19.43 7.107e-10 544-577
907	PR00988	URIDINE KINASE SIGNATURE	PR00988A 6.39 6.276e-12 314- 332
908	BL00107	Protein kinases ATP-binding region proteins.	BL00107A 18.39 5.950e-17 1125- 1156
909	BL00107	Protein kinases ATP-binding region proteins.	BL00107A 18.39 5.950e-17 1118- 1149
910	BL00107	Protein kinases ATP-binding region proteins.	BL00107A 18.39 8.560e-13 150- 181
911	BL00107	Protein kinases ATP-binding region proteins.	BL00107A 18.39 8.560e-13 150- 181
912	PF00856	SET domain proteins.	PF00856A 26.14 4.553e-11 243- 280
913	PF00628	PHD-finger.	PF00628 15.84 6.400e-13 197-212
914	PR00962	LETHAL(2) GIANT LARVAE PROTEIN SIGNATURE	PR00962D 10.40 1.000e-27 435- 459 PR00962G 15.71 4.086e-26 593-618 PR00962B 11.98 9.122e- 26 296-319 PR00962A 13.28 6.143e-22 15-34 PR00962C 8.00 4.000e-21 348-369 PR00962F 12.39 9.769e-21 552-572 PR00962H 13.32 2.636e-20 623- 643 PR00962I 11.68 9.786e-20 692-712 PR00962E 8.81 2.915e- 18 515-534
915	PR00962	LETHAL(2) GIANT LARVAE PROTEIN SIGNATURE	PR00962D 10.40 1.000e-27 365- 389 PR00962G 15.71 4.086e-26 523-548 PR00962A 13.28 6.143e- 22 15-34 PR00962C 8.00 4.000e- 21 278-299 PR00962F 12.39 9.769e-21 482-502 PR00962H

			1 0 1, 0 0 0 1, 0 10 7 0
SEO ACCESS	SION I.	DESCRIPTION	RESULTS*

SEQ IID NO:	ACCESSION NO.	· DESCRIPTION	RESULTS*
140.			13.32 2.636e-20 553-573 PR00962I 11.68 9.786e-20 622- 642 PR00962E 8.81 2.915e-18 445-464
916	BL00134	Serine proteases, trypsin family, histidine proteins.	BL00134A 11.96 5.886e-14 90- 107
917	BL00478	LIM domain proteins.	BL00478B 14.79 8.393e-13 211- 226 BL00478B 14.79 6.712e-10 271-286
918	PR00049	WILM'S TUMOUR PROTEIN SIGNATURE	PR00049D 0.00 5.729e-09 973- 988
922	BL00150	Acylphosphatase proteins.	BL00150 25.33 1.000e-40 37-84
924	DM00031	IMMUNOGLOBULIN V REGION.	DM00031B 15.41 8.063e-09 79- 113
925	BL00072	Acyl-CoA dehydrogenases proteins.	BL00072D 30.08 2.837e-24 280- 331 BL00072E 24.12 8.200e-24 368-411 BL00072C 25.30 7.873e- 20 226-267 BL00072B 9.48 6.049e-12 183-196
927	BL00237	G-protein coupled receptors proteins.	BL00237C 13.19 1.692e-13 229- 256 BL00237A 27.68 6.657e-13 90-130 BL00237D 11.23 9.571e- 13 290-307
928	BL01033	Globins profile.	BL01033A 16.94 7.923e-18 25-47 BL01033B 13.81 1.000e-15 93- 105
929	BL00216	Sugar transport proteins.	BL00216B 27.64 8.714e-13 203- 253
932	BL00415	Synapsins proteins.	BL00415N 4.29 9.519e-10 353- 397 BL00415N 4.29 2.117e-09 63-107 BL00415N 4.29 3.628e-09 57-101 BL00415N 4.29 5.664e-09 347-391
933	PD02448	TRANSCRIPTION PROTEIN DNA-BINDIN.	PD02448A 9.37 1.000e-40 46-85 PD02448B 10.17 1.000e-40 85- 133 PD02448C 13.62 1.000e-40 152-189 PD02448E 11.33 9.000e- 30 223-249 PD02448F 14.22 9.654e-25 267-291 PD02448D 11.48 3.659e-18 197-211 PD02448G 10.73 7.857e-16 293- 306
934	DM00191	w SPAC8A4.04C RESISTANCE SPAC8A4.05C DAUNORUBICIN.	DM00191D 13.94 9.083e-10 136- 175
935	BL01115	GTP-binding nuclear protein ran proteins.	BL01115A 10.22 4.696e-10 67- 111
936	BL00019	Actinin-type actin-binding domain proteins.	BL00019D 15.33 8.138e-14 865- 895
937	PR00762	CHLORIDE CHANNEL SIGNATURE	PR00762A 14.22 4.000e-22 183- 201 PR00762C 9.29 1.000e-21 268-288 PR00762E 12.07 3.250e- 20 520-537 PR00762D 11.29 1.000e-19 470-491 PR00762F 15.12 1.429e-19 538-558 PR00762B 12.12 1.818e-18 214- 234 PR00762G 14.13 3.455e-17 577-592
938	BL00027	'Homeobox' domain proteins.	BL00027 26.43 9.500e-25 291-334
939	DM01111	4 kw PHOSPHATASE	DM01111E 17.28 1.568e-10 248-

		DECORPORA	DECEM CO.
SEQ ID NO:	ACCESSION NO.	DESCRIPTION	RESULTS*
		TRANSFORMING 61K PDF1.	297 DM01111E 17.28 5.168e-10 659-708 DM01111D 16.76 5.263e-09 279-325 DM01111M 10.67 8.674e-09 911-935
940	BL00107	Protein kinases ATP-binding region proteins.	BL00107B 13.31 1.000e-14 293- 309 BL00107A 18.39 6.760e-13 229-260
942	BL01160	Kinesin light chain repeat proteins.	BL01160B 19.54 9.832e-11 543- 597
943	PD01066	PROTEIN ZINC FINGER ZINC- FINGER METAL-BINDING NU.	PD01066 19.43 3.500e-35 8-47
945	BL00989	Clathrin adaptor complexes small chain proteins.	BL00989B 26.51 1.000e-40 66- 117 BL00989A 11.66 1.000e-13 5-19
946	PR00178	FATTY ACID-BINDING PROTEIN SIGNATURE	PR00178D 13.52 9.571e-09 450- 469
947	BL00178	Aminoacyl-transfer RNA synthetases class-I proteins.	BL00178B 7.11 4.857e-09 713- 724
948	PF00628	PHD-finger.	PF00628 15.84 8.412e-14 201-216
951	BL00216	Sugar transport proteins.	BL00216B 27.64 2.050e-10 180- 230
952	PR00926	MITOCHONDRIAL CARRIER PROTEIN SIGNATURE	PR00926F 17.75 4.300e-11 26-49 PR00926F 17.75 6.348e-09 134- 157
955	PF00109	Beta-ketoacyl synthase.	PF00109 13.08 2.846e-12 342-357
957	PR00069	ALDO-KETO REDUCTASE SIGNATURE	PR00069A 16.01 8.826e-24 26-51 PR00069B 11.33 1.514e-17 86- 105 PR00069C 16.03 8.816e-14 155-173
958	PF00583	Acetyltransferase (GNAT) family.	PF00583A 12.53 5.500e-10 631- 642
961	PR00328	GTP-BINDING SAR1 PROTEIN SIGNATURE	PR00328A 10.62 8.740e-10 7-31
962	BL00354	HMG-I and HMG-Y DNA-binding domain proteins (A+T-hook).	BL00354A 3.83 9.438e-10 1489- 1499
963	BL00354	HMG-I and HMG-Y DNA-binding domain proteins (A+T-hook).	BL00354A 3.83 9.438e-10 1489- 1499
964	BL00027	'Homeobox' domain proteins.	BL00027 26.43 7.188e-27 53-96
965	PF00992	Troponin.	PF00992A 16.67 2.421e-09 581- 616
966	PR00515	5-HYDROXYTRYPTAMINE 1F RECEPTOR SIGNATURE	PR00515D 7.91 5.741e-09 13-33
967	BL00579	Ribosomal protein L29 proteins.	BL00579B 21.99 5.065e-21 164- 194
970	BL00504	Fumarate reductase / succinate dehydrogenase FAD-binding site proteins.	BL00504C 18.68 2.227e-24 34-59 BL00504D 10.43 7.261e-21 75-93
973	PF00580	UvrD/REP helicase.	PF00580A 13.37 4.720e-09 249- 271
974	PR00456	RIBOSOMAL PROTEIN P2 SIGNATURE	PR00456F 5.86 1.000e-10 242-254
975	BL00237	G-protein coupled receptors proteins.	BL00237A 27.68 4.429e-22 99- 139
976	BL00031	Nuclear hormones receptors DNA- binding region proteins.	BL00031A 19.55 7.158e-33 60-93 BL00031B 22.25 5.500e-28 94- 126
977	PD00066	PROTEIN ZINC-FINGER METAL- BINDI.	PD00066 13.92 8.200e-16 196-209 PD00066 13.92 8.200e-16 336-349 PD00066 13.92 2.385e-15 476-489

SEQ ID	ACCESSION NO.	DESCRIPTION	RESULTS*
NO:			77.0006610.000.000
			PD00066 13.92 9.308e-15 252-265
			PD00066 13.92 2.800e-14 448-461
1			PD00066 13.92 4.600e-14 392-405
			PD00066 13.92 5.200e-14 280-293
			PD00066 13.92 4.000e-13 224-237
			PD00066 13.92 4.429e-12 308-321
			PD00066 13.92 9.571e-12 420-433
			PD00066 13.92 6.870e-11 168-181
978	BL00721	Formatetetrahydrofolate ligase proteins.	BL00721B 13.21 1.000e-40 346-
			401 BL00721D 13.90 1.000e-40
			538-592 BL00721E 13.46 1.000e-
			40 597-646 BL00721I 18.79
			2.500e-40 814-860 BL00721H
			21.20 8.239e-39 763-814
			BL00721A 15.31 9.719e-32 287-
j)	321 BL00721C 16.92 4.000e-30
			498-535 BL00721F 15.96 8.232e-
			27 660-702 BL00721G 7.97
			3.017e-10 721-734
981	PD00126	PROTEIN REPEAT DOMAIN TPR	PD00126A 22.53 2.552e-09 180-
		NUCLEA.	201
982	BL00869	Renal dipeptidase proteins.	BL00869C 12.58 3.172e-19 59-95
			BL00869E 13.12 9.129e-18 120-
}			157 BL00869J 15.60 6.032e-17
			270-310 BL00869H 11.08 1.840e-
			16 219-242 BL00869G 13.55
			2.543e-16 192-214 BL00869F
			12.77 7.031e-14 157-192
			BL00869I 12.92 3.274e-12 242-
			270 BL00869D 14.02 5.282e-10
			95-124 BL00869B 15.55 9.382e-
			10 31-61
983	PR00196	ANNEXIN FAMILY SIGNATURE	PR00196F 13.89 2.125e-09 92-108
984	BL00485	Adenosine and AMP deaminase proteins.	BL00485D 30.82 2.427e-10 154-
		•	2.09

^{*} Results include in order: accession number subtype; raw score; p-value; position of signature in amino acid sequence

TABLE 4

SEQ ID NO:	PFAM NAME	DESCRIPTION	p-value	PFAM SCORE
2	ig	Immunoglobulin domain	3.9e-17	60.3
3	HSP90	Hsp90 protein	0	1548.4
6	tsp_1	Thrombospondin type 1 domain	0.002	22.1
7	7tm_1	7 transmembrane receptor (rhodopsin family)	6.7e-08	27.3
9	PWWP	PWWP domain	8.1e-16	66.0
12	Clq	Clq domain	1.7e-26	101.5
13	Clq	Clq domain	2e-20	81.3
14	Aa_trans	Transmembrane amino acid transporter protein	2.7e-42	153.9
15	E1-E2_ATPase	E1-E2 ATPase	6.3e-124	412.2
16	trypsin	Trypsin	1.2e-87	278.6
17	ig	Immunoglobulin domain	7.6e-12	43.2
18	lectin_c	Lectin C-type domain	0.0003	21.2
20	Alpha_L_fucos	Alpha-L-fucosidase	1.2e-217	736.5

11 0 02101220

SEQ ID NO:	PFAM NAME	DESCRIPTION	p-value	PFAM SCORE
22	pkinase	Eukaryotic protein kinase domain	3.3e-87	303.1
23	pkinase	Eukaryotic protein kinase domain	2.7e-85	296.8
24	pkinase	Eukaryotic protein kinase domain	2.7e-85	296.8
25	ank	Ank repeat	5.5e-14	59.9
27	pkinase	Eukaryotic protein kinase domain	1.5e-100	347.4
28	spectrin	Spectrin repeat	4e-57	203.2
29	spectrin	Spectrin repeat	4e-57	203.2
30	WD40	WD domain, G-beta repeat	1.2e-07	38.8
33	rrm	RNA recognition motif.	1.1e-17	72.2
34		RNA recognition motif.	1.1e-17	72.2
36	7tm_1	7 transmembrane receptor (rhodopsin family)	3e-36	117.3
37	ank	Ank repeat	5.9e-25	96.3
38	SRF-TF	SRF-type transcription factor	1.4e-36	133.9
40	alk phosphatase	Alkaline phosphatase	0	1034.9
	zf-C2H2	Zinc finger, C2H2 type	8.6e-103	354.9
44			3.1e-08	40.3
45	sugar_tr	Sugar (and other) transporter		275.6
47	7tm_2	7 transmembrane receptor (Secretin family)	6.4e-79	
50	zf-C2H2	Zinc finger, C2H2 type	1.3e-98	341.0
51	filament	Intermediate filament proteins	1.2e-176	600.3
52	zf-C3HC4	Zinc finger, C3HC4 type (RING finger)	2.7e-10	37.7
53	Cadherin_C_ter m	Cadherin cytoplasmic region	1.9e-94	327.2
54	S_100	S-100/ICaBP type calcium binding domain	5.2e-18	73.3
58	inositol_P	Inositol monophosphatase family	5e-13	49.8
59	7tm_1	7 transmembrane receptor (rhodopsin family)	8.8e-46	147.6
60	Kunitz_BPTI	Kunitz/Bovine pancreatic trypsin inhibito	3.7e-47	148.6
62	DAD	DAD family	2.5e-74	260.3
63	MOZ SAS	MOZ/SAS family	5.9e-133	455.1
64	MOZ_SAS	MOZ/SAS family	1.7e-123	423.6
65	ras	Ras family	9.3e-89	308.3
67	Ham lp like	Ham1 family	3.7e-49	176.7
68	7tm_l	7 transmembrane receptor (rhodopsin family)	5.2e-39	126.1
70	zf-C2H2	Zinc finger, C2H2 type	1.5e-112	387.3
71	Peptidase_M41	Peptidase family M41	1.2e-110	381.0
72	abhydrolase	alpha/beta hydrolase fold	9.8e-05	26.5
81	K tetra	K+ channel tetramerisation domain	0.022	-16.8
82		Eukaryotic protein kinase domain	5e-49	176.3
84	pkinase AAA	ATPases associated with various cellular act	1.3e-77	271.3
85	homeobox	Homeobox domain	1.4e-28	108.3
87	TGF-beta	Transforming growth factor beta like	6.7e-68	210.2
91		Mitochondrial carrier proteins	4.6e-57	198.5
95	mito_carr adenylatekinase	Adenylate kinase	1.1e-15	60.0
			4.1e-20	69.8
96	ig	Immunoglobulin domain		
99	CNH	CNH domain	3.4e-120	412.7
100	homeobox	Homeobox domain	7.4e-32	119.3
101	zf-C2H2	Zinc finger, C2H2 type	2.2e-47	170.8
102	zf-C2H2 ·	Zinc finger, C2H2 type	4.4e-89	309.4
103	dynamin	Dynamin family	1.4e-150	513.6
104	lectin_c	Lectin C-type domain	4.2e-15	63.6
105	lectin_c	Lectin C-type domain	4.2e-15	63.6
108	metalthio	Metallothionein	2e-25	97.9

SEQ ID NO:	PFAM NAME	DESCRIPTION	p-value	PFAM SCORE
112	HSP20	Hsp20/alpha crystallin family	2.6e-20	77.7
115	EF TS	Elongation factor TS	3.8e-63	221.1
116		Sugar (and other) transporter	4e-63	223.1
118	sugar_tr catalase	Catalase	0	1158.9
			1e-10	24.4
119	UCH	Ubiquitin carboxyl-terminal hydrolase, famil		
122	metalthio	Metallothionein	2.8e-25	97.4
125	adh_short	short chain dehydrogenase	1.6e-45	164.6
126	KRAB	KRAB box	7.9e-25	95.9
127	G-alpha	G-protein alpha subunit	1e-249	843.0
128	mito carr	Mitochondrial carrier proteins	2e-65	227.2
131	EF1BD	EF-1 guanine nucleotide exchange domain	4.9e-53	189.6
132	GYF	GYF domain	4.9e-28	106.6
133	GYF	GYF domain	4.9e-28	106.6
134	lipocalin	Lipocalin / cytosolic fatty-acid binding pr	2.1e-33	119.1
135	pkinase	Eukaryotic protein kinase domain	3.3e-86	299.8
136	ank	Ank repeat	2.2e-29	111.1
137	IL8	Small cytokines	3.1e-18	65.2
		(intecrine/chemokine), inter		
139	pyridoxal_deC	Pyridoxal-dependent decarboxylase conse	0.00011	19.0
140	cadherin	Cadherin domain	1.3e-88	307.8
142	efhand	EF hand	5.7e-33	123.0
143	Acyltransferase	Acyltransferase	2e-29	111.2
146	cytochrome_c	Cytochrome c	1.7e-33	124.7
147	pkinase	Eukaryotic protein kinase domain	2.3e-86	300.3
148	PDZ	PDZ domain (Also known as DHR or GLGF).	1.7e-09	45.0
149	aldo ket red	Aldo/keto reductase family	7.4e-189	640.8
150	homeobox	Homeobox domain	3.2e-08	38.7
151	PseudoU_synth_	tRNA pseudouridine synthase	4.7e-57	203.0
152	abhydrolase	alpha/beta hydrolase fold	1.7e-31	118.0
153	PDZ	PDZ domain (Also known as DHR or GLGF).	1.1e-09	45.6
156	PHD	PHD-finger	7.6e-15	62.8
157	fn3	Fibronectin type III domain	0.015	21.9
158	homeobox	Homeobox domain	2.7e-27	104.1
160	PWI	PWI domain	3.9e-24	93.6
162	DnaJ	DnaJ domain	2e-06	34.8
164	Cbl_N	CBL proto-oncogene N-terminal domain	8e-117	401.5
166	metalthio	Metallothionein	3.1e-26	100.6
167	LRR	Leucine Rich Repeat	0.00069	26.3
169	fibrinogen_C	Fibrinogen beta and gamma chains,	5.3e-180	611.4
170	fibrinogen_C	C-term Fibrinogen beta and gamma chains, C-term	5.3e-180	611.4
171	fibrinogen_C	Fibrinogen beta and gamma chains,	1e-149	510.8
100		C-term	1.6- 20	1116
173	homeobox	Homeobox domain	1.5e-29	111.6
174	FYVE	FYVE zinc finger	7.4e-28	103.8
175	GRIP	GRIP domain	3.9e-08	40.5
182	pkinase	Eukaryotic protein kinase domain	3.4e-71	250.0
185	CAP_GLY	CAP-Gly domain	5.6e-51	182.8
186	TBC	TBC domain	2.2e-50	180.8
187	TBC	TBC domain	2.2e-50	180.8

SEQ ID NO:	PFAM NAME	DESCRIPTION	p-value	PFAM SCORE
188	PDZ	PDZ domain (Also known as DHR or GLGF).	4e-13	57.0
189	Kelch	Kelch motif	5.2e-106	365.6
190	Tropomyosin	Tropomyosins	3.8e-171	535.4
192	Rieske	Rieske [2Fe-2S] domain	0.0016	18.5
199	ig	Immunoglobulin domain	5.9e-19	66.1
202	EGF	EGF-lik domain	3.4e-54	193.5
203 ·	trefoil	Trefoil (P-type) domain	1e-24	95.5
204	TBC	TBC domain	8.5e-38	139.0
205	efhand	EF hand	0.0096	22.6
206	ISK_Channel	Slow voltage-gated potassium channel	0.0031	8.1
207	trefoil	Trefoil (P-type) domain	2.9e-48	173.7
209	Ribosomal S13	Ribosomal protein S13/S18	1.2e-78	274.7
210	hemopexin	Hemopexin	1.3e-62	221.5
213	TBC \	TBC domain	2.5e-48	174.0
	Basic	Myogenic Basic domain	4.3e-50	179.8
215 216	Ribosomal L24	KOW motif	8.2e-23	89.2
	fn3	Fibronectin type III domain	7.3e-141	481.4
222	cofilin ADF	Cofilin/tropomyosin-type actin-	9.3e-47	168.8
223		binding pr		
224	efhand	EF hand	6.1e-06	33.2
225	Pterin_4a	Pterin 4 alpha carbinolamine dehydratase	9.3e-42	152.1
228	ABC_tran	ABC transporter	4.1e-110	379.2
234	E1_DerP2_DerF 2	E1 family	3.7e-90	312.9
235	E1_DerP2_DerF	E1 family	1.6e-48	174.6
237	PMP22 Claudin	PMP-22/EMP/MP20/Claudin family	1.7e-25	98.1
238	Opiods_neurope	Vertebrate endogenous opioids neurope	1.8e-159	543.2
239	eIF-5a	Eukaryotic initiation factor 5A hypusine	5.9e-104	358.8
240	Amino oxidase	Flavin containing amine oxidase	2.5e-11	37.8
243	zf-C2H2	Zinc finger, C2H2 type	2.1e-99	343.6
244	Band 7	SPFH domain / Band 7 family	2.3e-53	190.7
245	ank	Ank repeat	1.6e-88	307.5
246	zf-C2H2	Zinc finger, C2H2 type	6.7e-49	175.9
247	actin	Actin	2.3e-42	140.3
248	ER_lumen_recep	ER lumen protein retaining receptor	2.4e-155	529.5
250	PMP22_Claudin	PMP-22/EMP/MP20/Claudin family	2.2e-38	140.9
252	Collagen	Collagen triple helix repeat (20 copies)	1.4e-13	58.6
255	C2	C2 domain	0.052	7.8
257	CAP GLY	CAP-Gly domain	1.4e-20	81.8
260	WD40	WD domain, G-beta repeat	9.9e-62	218.5
261	WD40	WD domain, G-beta repeat	9.9e-62	218.5
262	WD40	WD domain, G-beta repeat	9.9e-62	218.5
263	cofilin_ADF	Cofilin/tropomyosin-type actin- binding pr	7.8e-21	82.6
264	Ribosomal L14	Ribosomal protein L14p/L23e	9.2e-10	40.6
7.044	<u> </u>	Saposin A-type domain	4.4e-27	103.4
	INAPA	I pobonit i e al bo continui		
265	SAPA	Sanosin A-type domain	4 4e-27	1 103.4
265 266	SAPA	Saposin A-type domain	4.4e-27	103.4
265 266 267	SAPA ABC_tran	ABC transporter	9.5e-39	142.2
265 266	SAPA		1	

SEQ ID NO:	PFAM NAME	DESCRIPTION	p-value	PFAM SCORE
273	rrm	RNA recognition motif.	0.074	14.6
275	lipocalin	Lipocalin / cytosolic fatty-acid binding pr	2.5e-41	146.4
276	ras	Ras family	1.1e-67	238.3
277	UCH	Ubiquitin carboxyl-terminal hydrolase, famil	1.2e-147	503.9
278	START	START domain	3.2e-09	44.1
279	WD40	WD domain, G-beta repeat	1.8e-27	104.7
282	G-patch	G-patch domain	7.8e-22	86.0
287	Anti_proliferat	BTG1 family	1.2e-101	351.0
289	KRAB	KRAB box	7.1e-21	82.8
293	7tm_3	7 transmembrane receptor	3.3e-73	256.6
295	SET	SET domain	5e-30	113.2
296	Pyridox_oxidase	Pyridoxamine 5'-phosphate oxidase	1.3e-76	268.0
297	rrm	RNA recognition motif.	5.4e-45	162.9
298	Ubie_methyltran	ubiE/COQ5 methyltransferase family	6.3e-05	-96.3
299	Ubie_methyltran	ubiE/COQ5 methyltransferase family	0.0024	-118.1
301	Cyt_reductase	FAD/NAD-binding Cytochrome reductase	7.7e-61	215.5
302	G-patch	G-patch domain	3.1e-14	60.7
307	7tm_1	7 transmembrane receptor (rhodopsin family)	7.7e-43	138.2
308	PH	PH domain	0.0015	17.8
310	7tm_1	7 transmembrane receptor (rhodopsin family)	1.4e-84	270.8
311	Rhodanese	Rhodanese-like domain	3.3e-64	226.7
312	tubulin	Tubulin/FtsZ family	4.9e-286	963.6
314	SURF4	SURF4 family	1.2e-199	676.6
325	IMS	impB/mucB/samB family	2e-58	207.5
327	cadherin	Cadherin domain	4.3e-91	316.0
329	NAC	NAC domain	2.1e-28	107.8
330	IP_trans	Phosphatidylinositol transfer protein	6.5e-98	338.7
332	TFIIS	Transcription factor S-II (TFIIS)	8.8e-05	29.3
337	zf-C2H2	Zinc finger, C2H2 type	3.6e-61	216.6
340	AIRS	AIR synthase related protein	4e-32 4.6e-80	120.2 279.4
343 346	annexin Stathmin	Annexin Stathmin family	1.8e-90	314.0
346	Ribosomal L16	Ribosomal protein L16	4.6e-09	34.9
348	lactamase B	Metallo-beta-lactamase superfamily	0.012	-6.0
351	efhand	EF hand	2.5e-14	61.0
353	lectin c	Lectin C-type domain	1.3e-05	32.1
354	WD40	WD domain, G-beta repeat	2.2e-18	74.5
360	lipocalin	Lipocalin / cytosolic fatty-acid binding pr	6.3e-10	38.3
362	Acetyltransf	Acetyltransferase (GNAT) family	0.0019	24.9
365	tRNA-synt_1	tRNA synthetases class I (I, L, M and V)	4.6e-185	628.2
366	Sulfatase	Sulfatase	6.1e-228	770.6
368	START	START domain	3.8e-11	50.5
369	pkinase	Eukaryotic protein kinase domain	2.4e-10	41.3
370	ACBP	Acyl CoA binding protein	4.4e-56	199.7
371	pkinase	Eukaryotic protein kinase domain	1.6e-94	327.5
373	EGF	EGF-like domain	2.6e-12	54.3
375	zf-C2H2	Zinc finger, C2H2 type	8.2e-64	225.4
377	KRAB	KRAB box	3.7e-27	103.7
379	SET	SET domain	7.3e-61	215.6
380	Glyc _transf_8	Glycosyl transferase family 8	0.0028	-40.1
381	zf-C2H2	Zinc finger, C2H2 type	4.3e-06	33.7
383	Glyco_transf_8	Glycosyl transferase family 8	0.0028	-40.1

SEQ ID NO:	PFAM NAME	DESCRIPTION	p-value	PFAM SCORE
384	RasGEF	RasGEF domain	8.1e-43	155.7
385	TBC	TBC domain	0.017	-66.6
389	Glycos transf 2	Glycosyl transferases	1.3e-15	65.3
390	Na Ca Ex	Sodium/calcium exchanger protein	3.9e-105	362.7
391	fn3	Fibronectin type III domain	4.1e-102	352.6
392	fn3	Fibronectin type III domain	3.4e-45	163.6
393	fn3	Fibronectin type III domain	3.4e-45	163.6
394	ldl_recept_b	Low-density lipoprotein receptor	7.1e-49	175.8
395	Ribosomal L30	repeat Ribosomal protein L30p/L7e	0.0023	16.0
396	Oxysterol BP	Oxysterol-binding protein	1.5e-94	327.5
397	RDS ROM1	Peripherin/rom-1	2.9e-33	123.9
399	lactamase B	Metallo-beta-lactamase superfamily	3.4e-39	143.6
402	F-box	F-box domain.	0.0002	28.1
403	CLP_protease	Clp protease	4.8e-64	226.2
405	Ribosomal_L35	Ribosomal protein L35Ae	6e-77	269.0
406	LIM	LIM domain containing proteins	0.00021	20.7
410	tRNA-synt 1c	tRNA synthetases class I (E and Q)	1e-236	799.8
411	NTP transf 2	Nucleotidyltransferase domain	3.9e-16	67.0
412	DEAD	DEAD/DEAH box helicase	0.00016	17.2
414	DUF94	Domain of unknown function DUF94	0.00011	26.9
415	tubulin	Tubulin/FtsZ family	4.5e-289	973.7
420	SET	SET domain	3.3e-57	203.5
421	WD40	WD domain, G-beta repeat	6.1e-29	109.6
423	zf-C2H2	Zinc finger, C2H2 type	1.5e-39	144.9
424	pkinase	Eukaryotic protein kinase domain	8.9e-75	261.8
428	LIM	LIM domain containing proteins	1.8e-34	126.7
431	kazal	Kazal-type serine protease inhibitor domain	3.7e-18	73.8
432	SH2	Src homology domain 2	1.4e-67	198.4
433	zf-C2H2	Zinc finger, C2H2 type	2.8e-144	492.7
434	ras	Ras family	0.012	-106.8
436	E1-E2 ATPase	E1-E2 ATPase	1.6e-117	391.0
437	RNA pol A	RNA polymerase alpha subunit	0	1077.7
438	PHD	PHD-finger	1.6e-11	51.7
439	lectin c	Lectin C-type domain	4.7e-30	113.3
440	zf-C2H2	Zinc finger, C2H2 type	1.1e-65	231.6
441	arrestin	Arrestin (or S-antigen)	2.9e-254	858.1
442	aminotran 3	Aminotransferases class-III	8.2e-80	231.1
	_	pyridoxal-pho		
443	UCH-1	Ubiquitin carboxyl-terminal hydrolases famil	8.5e-12	52.6
444	CTF_NFI	CTF/NF-I family	2.6e-277	934.6
451	T-box	T-box	3.8e-117	402.6
453	Rieske	Rieske [2Fe-2S] domain	2.6e-13	57.7
454	zf-C2H2	Zinc finger, C2H2 type	3.9e-64	226.5
456	homeobox	Homeobox domain	2.8e-08	38.9
459	ig	Immunoglobulin domain	2.6e-20	70.5
460	Hydrolase	haloacid dehalogenase-like hydrolase	4e-25	96.9
462	rve	Integrase core domain	1.6e-13	50.7
466	CH	Calponin homology (CH) domain	2.4e-17	71.1
467	CH	Calponin homology (CH) domain	2.4e-17	71.1
468	Sterol desat	Sterol desaturase	7.5e-38	139.2
469	pro_isomerase	Cyclophilin type peptidyl-prolyl cis- tr	2.6e-63	220.9
470	Peptidase M24	metallopeptidase family M24	6e-08	28.1
471	PDZ	PDZ domain (Also known as DHR or GLGF).	5.4e-129	441.9

SEQ ID	PFAM NAME	DESCRIPTION	p-value	PFAM
NO:				SCORE
472	myb_DNA- binding	Myb-like DNA-binding domain	3.6e-06	33.9
473	ZZ	Zinc finger present in dystrophin, CB	0.012	20.0
474	EF1G_domain	Elongation factor 1 gamma, conserved doma	6.3e-88	305.5
475	Ribosomal_L31e	Ribosomal protein L31e	6.1e-66	232.5
476	Clq	C1q domain	2.5e-75	263.7
477	SH3	SH3 domain	1.1e-12	55.6
478	MoaA_NifB_Pq qE	moaA / nifB / pqqE family	0.002	-17.7
479	FYVE	FYVE zinc finger	9.3e-21	78.6
480	DNA_pol_A	DNA polymerase family A	2.3e-46	167.4
482	adh_short	short chain dehydrogenase	1.2e-62	221.6
483	ank	Ank repeat	1.3e-17	71.9
484	IMS	impB/mucB/samB family	2.2e-83	290.5
486	TIR	TIR domain	3.2e-19	67.8
487	FMO-like	Flavin-binding monooxygenase-like	0	1425.5
488	I_LWEQ	I/LWEQ domain	9.5e-101	341.0
495	homeobox	Homeobox domain	3.6e-06	30.8
497	pkinase	Eukaryotic protein kinase domain	2.3e-166	566.1
499	fn3	Fibronectin type III domain	2.5e-237	801.8 115.6
501	LRR	Leucine Rich Repeat Regulator of G protein signaling	9.3e-31 0.041	115.6
502	RGS	domain		
503	filament	Intermediate filament proteins	1e-142	487.5
505	fn3	Fibronectin type III domain	1.3e-100	347.7
506	HECT	HECT-domain (ubiquitin- transferase).	1e-13	59.0
507	Ribosomal_L7A e	Ribosomal protein L7Ae	5.7e-26	99.7
508	WD40	WD domain, G-beta repeat	0.063	19.8
509	WD40	WD domain, G-beta repeat	0.063	19.8
510	WD40	WD domain, G-beta repeat	2.1e-42	154.3
511	pkinase	Eukaryotic protein kinase domain	2.3e-86	300.4
512	G-gamma	GGL domain	1.9e-08	34.3
513	SH3	SH3 domain	3e-06	34.2
515	HTH_AraC	Bacterial regulatory helix-turn-helix protei	3.9e-27	103.6
516	zf-C2H2	Zinc finger, C2H2 type	1.7e-34	128.0
517	S1	S1 RNA binding domain	6.1e-58	205.9
518	pkinase	Eukaryotic protein kinase domain	1.8e-75	264.2
525	cadherin	Cadherin domain	2e-80	280.6
528	zf-C2H2	Zinc finger, C2H2 type	4e-70	246.4
529	neur_chan	Neurotransmitter-gated ion-channel	5.8e-222	750.8
531	RhoGEF	RhoGEF domain	3.5e-44	160.2
532	myosin_head	Myosin head (motor domain)	0	1494.5
533	LRR	Leucine Rich Repeat	8.3e-15	62.6
535	Sec7	Sec7 domain	5.1e-92 4.8e-05	319.1 26.4
536	homeobox	Homeobox domain	4.8e-05 2.4e-100	330.6
539	actin	Ank repeat	1.9e-35	131.2
542	ank zf-CCCH	Ank repeat	1.9e-35 2.8e-10	41.7
544 546	DSPc	Zinc finger C-x8-C-x5-C-x3-H type Dual specificity phosphatase,	2.8e-10 2.4e-40	147.4
		catalytic doma		
547	HMG_CoA_synt	Hydroxymethylglutaryl-coenzyme A synthas	0	1250.8
549	laminin_G	Laminin G domain	3.3e-76	266.6
			1 0 000	102
551 552	PHD	PHD-finger PDZ domain (Also known as DHR or	0.008	9.3 25.0

SEQ ID NO:	PFAM NAME	DESCRIPTION	p-value	PFAM SCORE
		GLGF).		
555	WW	WW domain	1.3e-24	95.3
558	kinesin	Kinesin motor domain	1.8e-176	599.7
559	zf-C3HC4	Zinc finger, C3HC4 type (RING finger)	0.00085	16.5
563	efhand	EF hand	7.9e-11	49.4
567	PH	PH domain	7.8e-06	25.9
568	PH	PH domain	3.1e-39	143.8
569	Hist_deacetyl	Histone deacetylase family	5.2e-106	365.6
570	PDZ	PDZ domain (Also known as DHR or GLGF).	3.4e-20	80.5
571	zf-C3HC4	Zinc finger, C3HC4 type (RING finger)	le-16	58.5
573	ubiquitin	Ubiquitin family	1.4e-08	31.1
574	FH2	Formin Homology 2 Domain	1.3e-110	380.9
576	serpin	Serpins (serine protease inhibitors)	4.3e-146	496.4
579	zf-C2H2	Zinc finger, C2H2 type	5.7e-76	265.8
580	pkinase	Eukaryotic protein kinase domain	6.9e-79	275.5
581	RhoGAP	RhoGAP domain	4.4e-53	189.8
582	Ribosomal_L7A	Ribosomal protein L7Ae	0.028	1.0
584	kazal	Kazal-type serine protease inhibitor domain	2.2e-52	187.4
585	LRR	Leucine Rich Repeat	4.4e-28	106.7
586	PHD	PHD-finger	3.8e-12	53.8
588	GTP1_OBG	GTP1/OBG family	1.1e-62	215.2
590	Collagen	Collagen triple helix repeat (20 copies)	8e-42	152.4
591	lys .	C-type lysozyme/alpha-lactalbumin family	1.6e-31	116.4
596 .	ACBP	Acyl CoA binding protein	0.0022	-9.4
597	SNF2_N	SNF2 and others N-terminal domain	3.7e-98	339.5
600	KRAB	KRAB box	1.3e-29	111.8
606	LRR	Leucine Rich Repeat	1e-05	32.5
607	LRR	Leucine Rich Repeat	1e-05	32.5
608	WD40	WD domain, G-beta repeat	5.3e-23	89.8
610	cpn60_TCP1	TCP-1/cpn60 chaperonin family	1.7e-237	802.4
613	THF_DHG_CY H	Tetrahydrofolate dehydrogenase/cyclohydro	4.9e-173	588.3
617	rrm	RNA recognition motif.	4e-14	60.4
618	rrm	RNA recognition motif.	4e-14	60.4
620	cofilin_ADF	Cofilin/tropomyosin-type actin- binding pr	3e-06	34.2
621	Nop	Putative snoRNA binding domain	6.1e-95	328.8
622	UCH-2	Ubiquitin carboxyl-terminal hydrolase family	5.8e-21	83.1
625	zf-C2H2	Zinc finger, C2H2 type	2.5e-124	426.4
628	DEAD	DEAD/DEAH box helicase	2.5e-68	219.0
632	GST	Glutathione S-transferases.	4.8e-26	89.0
633	5 nucleotidase	5'-nucleotidase	6.6e-248	837.0
636	LIM	LIM domain containing proteins	1.6e-88	307.5
637	pkinase	Eukaryotic protein kinase domain	1.5e-73	257.8
638	MSP domain	MSP (Major sperm protein) domain	8.4e-09	42.7
639	metalthio	Metallothionein	2e-24	94.6
641	zf-C2H2	Zinc finger, C2H2 type	6.1e-114	391.9
642	Ribosomal S28e	Ribosomal protein S28e	9.3e-48	172.1
643	Ribosomal S5	Ribosomal protein S5	8.3e-87	301.8
646	PHD	PHD-finger	0.00025	23.1

SEQ ID NO:	PFAM NAME	DESCRIPTION	p-value	PFAM SCORE
648	Lipase_GDSL	Lipase/Acylhydrolase with GDSL- like motif	0.015	2.2
652	zf-C2H2	Zinc finger, C2H2 type	4.1e-146	498.8
653	histone	Core histone H2A/H2B/H3/H4	1.2e-10	48.8
654	zf-C2H2	Zinc finger, C2H2 type	1.9e-87	303.9
655	ras	Ras family	6.4e-77	269.0
657	zf-C3HC4	Zinc finger, C3HC4 type (RING finger)	5.3e-13	46.4
658	STphosphatase	Ser/Thr protein phosphatase	2.6e-182	619.1
659	zf-C2H2	Zinc finger, C2H2 type	1.3e-92	321.1
660	zf-C2H2	Zinc finger, C2H2 type	1.5e-85	297.6
662	NDK	Nucleoside diphosphate kinases	1.4e-119	410.7
664	IRF	Interferon regulatory factor transcription f	7e-20	79.5
665	4HPPD_C	4-hydroxyphenylpyruvate dioxygenase C term	1.4e-16	68.5
666	DEAD	DEAD/DEAH box helicase	4.8e-74	237.1
667	DEAD	DEAD/DEAH box helicase	2.9e-70	225.1
669	pkinase	Eukaryotic protein kinase domain	6.1e-93	322.2
671	homeobox	Homeobox domain	0.018	16.5
678	crystall	Beta/Gamma crystallin	4.7e-106	365.8
679	WD40	WD domain, G-beta repeat	1.9e-06	34.9
680	Keratin B2	Keratin, high sulfur B2 protein	4.1e-06	15.9
682	G-gamma	GGL domain	8.5e-33	117.9
685	UCH-2	Ubiquitin carboxyl-terminal hydrolase family	1.4e-29	111.7
686	Acetyltransf	Acetyltransferase (GNAT) family	6.6e-10	46.4
687	7tm_1	7 transmembrane receptor (rhodopsin family)	4.6e-15	50.0
688	proteasome	Proteasome A-type and B-type	6.5e-64	225.7
689	SCP2	SCP-2 sterol transfer family	6.2e-37	136.1
690	TS-N	TS-N domain	0.041	20.1
692	zf-C2H2	Zinc finger, C2H2 type	9.9e-60	211.9
693	zf-MYND	MYND finger	0.038	5.5
694	Oxysterol BP	Oxysterol-binding protein	3.9e-133	455.7
695	PDZ	PDZ domain (Also known as DHR or GLGF).	1.3e-30	115.1
703	Peptidase C2	Calpain family cysteine protease	2.3e-175	596.0
706	filament	Intermediate filament proteins	7.2e-107	368.5
710	fibrinogen_C	Fibrinogen beta and gamma chains, C-term	7e-80	278.0
711 ·	SH2	Src homology domain 2	2.3e-65	192.1
712	ATP-synt_DE	ATP synthase, Delta/Epsilon chain	0.00062	19.0
713	ARID	ARID DNA binding domain	2e-17	71.3
714	LBP BPI CETP	LBP / BPI / CETP family	8.6e-34	125.7
715	RNA_pol_L	RNA polymerases L / 13 to 16 kDa subunit	4.8e-49	176.3
716	KRAB	KRAB box	1.3e-42	155.0
717	mito_carr	Mitochondrial carrier proteins	4.8e-38	133.3
719	Gal-bind lectin	Vertebrate galactoside-binding lectin	1.5e-25	90.2
726	aldedh	Aldehyde dehydrogenase family	1.3e-119	410.8
728	Glycos transf 2	Glycosyl transferases	4e-21	83.6
734	ELM2	ELM2 domain	2e-34	127.8
735	PR55	Protein phosphatase 2A regulatory subunit PR	0	1038.2
737	DSPc	Dual specificity phosphatase, catalytic doma	4e-14	60.4
740	WD40	WD domain, G-beta repeat	5.6e-14	59.9
745	zf-C3HC4	Zinc finger, C3HC4 type (RING	3.8e-13	46.9

SEQ ID PFAM NAME NO:		DESCRIPTION	p-value	PFAM SCORE
110.		finger)		
749	mito carr	Mitochondrial carrier proteins	4.5e-67	232.8
750	DUF27	Domain of unknown function DUF27	4.5e-12	53.5
751	SH3	SH3 domain	3.6e-17	70.5
752	HMG box	HMG (high mobility group) box	8.6e-13	55.9
753	SPRY	SPRY domain	5.9e-05	23.3
754	GTP CDC	Cell division protein	7.5e-153	521.2
755	mito carr	Mitochondrial carrier proteins	3e-88	305.4
756	TSPN	Thrombospondin N-terminal -like	8.1e-58	205.5
		domains		
757	BTB	BTB/POZ domain	5.7e-23	89.7
759	zf-C2H2	Zinc finger, C2H2 type	1.2e-12	55.4
760	NSF	NSF attachment protein	6.4e-127	435.1
762	Ribosomal_S14	Ribosomal protein S14p/S29e	2.1e-06	24.8
765	ThiF_family	ThiF family	1.7e-39	144.6
766	DnaJ	DnaJ domain	3.9e-36	133.5
768	tRNA-synt_2b	tRNA synthetase class II	9.1e-81	281.7
769	ldl_recept_a	Low-density lipoprotein receptor domain	0	1404.5
770	WD40	WD domain, G-beta repeat	2e-21	84.6
771	LRR	Leucine Rich Repeat	3.8e-06	33.9
774	SNF2 N	SNF2 and others N-terminal domain	5.5e-99	342.3
776	VPS9	Vacuolar sorting protein 9 (VPS9)	1.1e-30	115.4
	1 1 1 1	domain		1
777	VPS9	Vacuolar sorting protein 9 (VPS9) domain	1.1e-30	115.4
778	VPS9	Vacuolar sorting protein 9 (VPS9) domain	1.1e-30	115.4
779	zf-C3HC4	Zinc finger, C3HC4 type (RING finger)	3.1e-08	31.0
781	cadherin	Cadherin domain	5.6e-113	388.7
783	HECT	HECT-domain (ubiquitin-	4.2e-31	116.8
		transferase).		
785	sushi	Sushi domain (SCR repeat)	1.8e-60	214.3
786	sushi	Sushi domain (SCR repeat)	1.8e-60	214.3
788	vwa	von Willebrand factor type A domain	1.9e-52	187.7
790	rrm	RNA recognition motif.	2.8e-20	80.8
791	Collagen	Collagen triple helix repeat (20 copies)	0.00097	9.7
792	pkinase	Eukaryotic protein kinase domain	0.023	12.4
795	zf-C2H2	Zinc finger, C2H2 type	6.5e-95	328.7
796	adh short	short chain dehydrogenase	4.1e-05	-7.3
799	SAICAR synt	SAICAR synthetase	6e-125	428.5
805	WD40	WD domain, G-beta repeat	4e-65	229.8
806	ZU5	ZU5 domain	4.7e-37	136.5
807	WD40	WD domain, G-beta repeat	0.016	21.8
808	WD40	WD domain, G-beta repeat	0.0041	23.8
809	pkinase	Eukaryotic protein kinase domain	2e-31	117.2
810	vwa	von Willebrand factor type A domain	1.9e-52	187.7
814	zf-C2H2	Zinc finger, C2H2 type	4.5e-83	289.4
815	zf-C2H2	Zinc finger, C2H2 type Zinc finger, C2H2 type	6e-74	259.1
			l	
817	myosin_head	Myosin head (motor domain)	1.5e-176	599.9
818	GSPII_E	Bacterial type II secretion system protein	0.012	11.5
819	PDEase	3'5'-cyclic nucleotide phosphodiesterase	1.1e-74	215.5
821	PH	PH domain	0.00025	20.5
822	CNH	CNH domain	0.00015	-24.7
		RNA recognition motif.	1.5e-06	1

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SEQ ID NO:	PFAM NAME	DESCRIPTION	p-value	PFAM SCORE
829	HMG box	HMG (high mobility group) box	7.8e-34	125.8
830	RasGEF	RasGEF domain	2.2e-102	353.5
831	CNH	CNH domain	3e-118	406.2
832	mito carr	Mitochondrial carrier proteins	3.7e-37	130.3
833	PX	PX domain	2.7e-19	77.5
837	Y phosphatase	Protein-tyrosine phosphatase	1.6e-263	888.8
838	ank	Ank repeat	2.4e-270	911.5
840	ank	Ank repeat	5.8e-38	139.6
842	Ribosomal L15e	Ribosomal L15	4.8e-131	448.8
843	SNF	Sodium:neurotransmitter symporter family	0	1201.8
845	Peptidase M16	Insulinase (Peptidase family M16)	4.7e-67	236.2
848	EF1BD	EF-1 guanine nucleotide exchange	2.2e-56	200.7
•		domain		
849	zf-C2H2	Zinc finger, C2H2 type	1.5e-122	420.5
850	zf-C2H2	Zinc finger, C2H2 type	2e-67	237.4
852	SIS	SIS domain	3.8e-30	113.6
853	RhoGAP	RhoGAP domain	1.1e-37	138.6
854	PDZ	PDZ domain (Also known as DHR or	5.1e-10	46.7
056	ACOY	GLGF).	9.1e-263	886.3
856	ACOX	Acyl-CoA oxidase		
858	efhand	EF hand	2.4e-18	74.4
860 .	homeobox	Homeobox domain	4e-22	86.9
862	TFIIF_beta	Transcription initiation factor IIF, beta	2.2e-134	459.8
866	A2M	Alpha-2-macroglobulin family	4.9e-21	70.9
867	MoCF_biosynth	Molybdenum cofactor biosynthesis protei	5.8e-205	694.3
868	EGF	EGF-like domain	4.1e-22	86.9
869	EGF	EGF-like domain	1.1e-22	88.8
871	PI-PLC-X	Phosphatidylinositol-specific phospholipase	7.2e-95	328.6
872	UCH-2	Ubiquitin carboxyl-terminal hydrolase family	1.1e-20	82.1
874	SH3	SH3 domain	2.2e-14	61.2
877	SH3	SH3 domain	8.6e-90	311.7
882	KRAB	KRAB box	6.9e-45	162.6
885	ank	Ank repeat	7.1e-07	36.3
886	biopterin_H	Biopterin-dependent aromatic amino acid h	0	988.3
887	GTP EFTU	Elongation factor Tu family	4.9e-129	437.5
888	zf-C3HC4	Zinc finger, C3HC4 type (RING finger)	1.6e-14	51.4
889	zf-C2H2	Zinc finger, C2H2 type	3.7e-92	319.6
890	ig	Immunoglobulin domain	3.8e-06	24.8
892	PTR2	POT family	9.5e-48	163.0
893	Sulfatase	Sulfatase	3.5e-78	273.2
894	Sulfatase	Sulfatase	3.5e-78	273.2
895	7tm_1	7 transmembrane receptor (rhodopsin family)	4.5e-51	164.4
896	Glyco_hydro_31	Glycosyl hydrolases family 31	0	1277.3
897	chromo	'chromo' (CHRromatin Organization MOdifier)	3.9e-06	26.0
898	Cbl_N	CBL proto-oncogene N-terminal domain	1.2e-273	922.4
899	vwa	von Willebrand factor type A domain	5.5e-32	119.7
900	WD40	WD domain, G-beta repeat	2.7e-07	37.7
901	zf-C2H2	Zinc finger, C2H2 type	4e-156	532.1
903	ras	Ras family	6.6e-101	348.6

SEQ ID NO:	PFAM NAME	DESCRIPTION	p-value	PFAM SCORE
904	Armadillo_seg	Armadillo/beta-catenin-like repeats	1.1e-06	35.6
906	FH2	Formin Homology 2 Domain	4.5e-112	385.7
907	Cytidylyltransf	Cytidylyltransferase	1.4e-05	29.3
908	pkinase	Eukaryotic protein kinase domain	1.2e-64	228.2
909	pkinase	Eukaryotic protein kinase domain	8.5e-70	245.3
910	pkinase	Eukaryotic protein kinase domain	2.9e-42	153.8
911	pkinase	Eukaryotic protein kinase domain	1.2e-35	131.8
912	PHD	PHD-finger	5.1e-06	33.4
913	PHD	PHD-finger	5.5e-16	66.5
916	filament	Intermediate filament proteins	9.7e-121	414.5
917	LIM	LIM domain containing proteins	5.9e-15	57.9
918	SAM	SAM domain (Sterile alpha motif)	4.3e-16	66.9
922	Acylphosphatase	Acylphosphatase	2.9e-63	223.6
924	ig	Immunoglobulin domain	1.3e-08	32.8
925	Acyl-CoA_dh	Acyl-CoA dehydrogenase	2.4e-131	449.8
927	7tm_1	7 transmembrane receptor (rhodopsin family)	2.9e-45	145.9
928	globin	Globin	2.4e-52	186.9
929	sugar_tr	Sugar (and other) transporter	1.2e-16	68.8
932	Collagen	Collagen triple helix repeat (20 copies)	0.00097	9.7
933	HMG_box	HMG (high mobility group) box	7.8e-34	125.8
934	SEA	SEA domain	0.0021	24.7
935	ras	Ras family	6.4e-59	209.2
936	CH	Calponin homology (CH) domain	3.8e-21	83.7
937	voltage_CLC	Voltage gated chloride channels	1.9e-199	676.0
938	homeobox	Homeobox domain	1.9e-25	98.0
940	pkinase	Eukaryotic protein kinase domain	9.9e-58	205.2
942	Myosin_tail	Myosin tail	3.7e-09	38.2
943	zf-C2H2	Zinc finger, C2H2 type	2.2e-92	320.3
945	Clat_adaptor_s	Clathrin adaptor complex small chain	1.3e-76	268.0
946	sugar_tr	Sugar (and other) transporter	0.017	-122.8
947	tRNA-synt_le	tRNA synthetases class I (C)	0.00097	15.6
948	PHD	PHD-finger	2.2e-17	71.2
951	sugar_tr	Sugar (and other) transporter	0.0082	-113.9
952	mito_carr	Mitochondrial carrier proteins	1.7e-54	189.7
953	myb_DNA- binding	Myb-like DNA-binding domain	4.5e-20	80.1
955	ketoacyl-synt	Beta-ketoacyl synthase	7.1e-133	454.8
957	aldo_ket_red	Aldo/keto reductase family	1.5e-98	340.8
959	Kelch	Kelch motif	0.02	20.8
961	ras	Ras family	2.2e-29	111.1
964	homeobox	Homeobox domain	5.4e-22	86.5
965	PH	PH domain	3e-21	80.9
966	zf-C3HC4	Zinc finger, C3HC4 type (RING finger)	2.2e-09	34.7
967	Ribosomal_L29	Ribosomal L29 protein	1.6e-15	65.0
970	FAD_binding_2	FAD binding domain	8.9e-47	166.6
971	rve	Integrase core domain	0.00015	19.8
972	Glycos_transf_2	Glycosyl transferases	2.1e-21	84.5
974	Ribosomal_L10	Ribosomal protein L10	3.3e-48	173.6
975	7tm_1	7 transmembrane receptor (rhodopsin family)	1.6e-37	121.3
976	zf-C4 .	Zinc finger, C4 type (two domains)	2.1e-52	178.5
977	zf-C2H2	Zinc finger, C2H2 type	6.6e-150	511.4
978	FTHFS	Formate-tetrahydrofolate ligase	0	1367.2
982	Renal_dipeptase	Renal dipeptidase	1.3e-73	258.0
984	A deaminase	Adenosine/AMP deaminase	2.6e-05	-48.6

SEQ ID NO: of full-length nucleotide sequence	SEQ ID NO: of full-length peptide	SEQ ID NO: of contig nucleotide sequence	SEQ ID NO: f contig peptide sequence	Priority docket number_corresp ndin g SEQ ID NO: in priority application	SEQ ID NO: in U.S.S.N. 09/496,914
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4	988	1972	2956	787CIP2_4	2123
5	989	1973	2957	787CIP2_5	
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7	991	1975	2959	787CIP2_7	3324
8	992	1976	2960	787CIP2_8	6182
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11	995	1979	2963	787CIP2_11	6257
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44	1028	2012	2996	787CIP2 44	7022
45	1029	2013	2997	787CIP2 46	7057
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49	1032	2017	3001	787CIP2_51	7182
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93	1077	2061	3045	787CIP2 95	7914
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103	1087	2071	3055	787CIP2_105	7944
104	1088	2072	3056	787CIP2_106	7951
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109	1093	2077	3061	787CIP2_111	7978
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112	1096	2080	3064	787CIP2_114	8000
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115	1099	2083	3067	787CIP2_117	8007
116	1100	2084	3068	787CIP2_118	8008
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119	1103	2087	3071	787CIP2_121	8017
120	1104	2088	3072	787CIP2_122	8018
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131	1115	2099	3083	787CIP2_133	8045
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135	1119	2103	3087	787CIP2_137	8053
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151	1135	2119	3103	787CIP2 153	8087
152	1136	2120	3104	787CIP2 154	8091
153	1137	2121	3105	787CIP2 155	8100
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171	1155	2139	3123	787CIP2_173	8140
172	1156	2140	3124	787CIP2_174	8141
173	1157	2141	3125	787CIP2_175	8147

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294	1278	2262	3246	787CIP2 299	10335
295	1279	2263	3247	787CIP2 300	10336
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305	1289	2273	3257	787CIP2_311	10348
306	1290	2274	3258	787CIP2_312	10349
307	1291	2275	3259	787CIP2_314	10351
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309	1293	2277	3261	787CIP2_316	10353
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312	1296	2280	3264	787CIP2_319	10356
313	1297	2281	3265	787CIP2_320	10357
314	1298	2282	3266	787CIP2_321	10358
315	1299	2283	3267	787CIP2_322	10360
316	1300	2284	3268	787CIP2_323	10361
317	1301	2285	3269	787CIP2_324	10362
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319	1303	2287	3271	787CIP2_326	10365
320	1304	2288	3272	787CIP2_327	10366
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338	1322	2306	3290	787CIP2_348	10393
339	1323	2307	3291	787CIP2_349	10394
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F-414	11200	1 2202	1 2266	Leagornon (a	1 (017
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425	1410	2394	3378	787CIP2B_74	6269
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456	1440	2424	3408	787CIP2B 105	6425
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538	1522	2507	3490	787CIP2B_187	6849
539	1523	2508	3491	787CIP2B_188	6851
540	1524		3493	787CIP2B_189	6851
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542	1526	2510 2511	3494	787CIP2B_191 787CIP2B 192	6869
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544 545	1528 1529	2513	3497	787CIP2B 194	6887
546	1530	2514	3498	787CIP2B 195	6890
547	1531	2515	3499	787CIP2B 196	6894
548	1532	2516	3500	787CIP2B 197	6899
		2517	3501	787CIP2B 198	6900
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		2519	3503	787CIP2B 200	6910
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576	1560	2544	3528	787CIP2B_226	7016
577	1561	2545	3529	787CIP2B_227	7023
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582	1566	2550	3534	787CIP2B_232	7040
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591	1575	2559	3543	787CIP2B_241	7079
592	1576	2560	3544	787CIP2B_242	7085
593	1577	2561	3545	787CIP2B_243	7148

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		0.500	T 2016	Lagacinon C44	7156
594	1578	2562	3546	787CIP2B_244	7156 .
595	1579	2563	3547	787CIP2B_245	7171
596	1580	2564	3548	787CIP2B_246 787CIP2B_248	7265
597	1581	2565	3549 3550	787CIP2B_248 787CIP2B_249	7268
598	1582	2566	3551	787CIP2B_249 787CIP2B_250	7308
599	1583	2567	3552	787CIP2B_230 787CIP2B_251	7336
600	1584	2568	3553	787CIP2B_231	7347
601	1585	2569	3554	787CIP2B_252 787CIP2B 253	7405
602	1586	2570	3555	787CIP2B_253	7405
603	1587	2571	3556	787CIP2B_254	7403
604	1588	2572 2573	3557	787CIP2B_255	7412
605	1589	2574	3558	787CIP2B_250	7436
606	1590	2575	3559	787CIP2B 258	7436
607	1591	2576	3560	787CIP2B 259	7454
608	1592		3561	787CIP2B 260	7476
609	1593	2577	3562	787CIP2B_200	7598
610	1594	2578 2579		787CIP2B 262	7619
611	1595	2579	3563 3564	787CIP2B_202 787CIP2B_263	7644
612	1596		3565	787CIP2B_263	7648
613	1597	2581 2582	3566	787CIP2B_265	7659
614	1598	2582	3567	787CIP2B_265	7661
615	1599		3568	787CIP2B 267	7669
616	1600	2584	3569	787CIP2B_267	7686
617	1601	2585 2586	3570	787CIP2B_268	7686
618	1602	2587	3571	787CIP2B 270	7694
619		2588	3572	787CIP2B 271	7697
620 621	1604 1605	2589	3573	787CIP2B_271	7733
621	1605	2590	3574	787CIP2B_272	7734
623	1607	2591	3575	787CIP2B 274	7744
624	1607	2592	3576	787CIP2B 275	7751
625	1609	2593	3577	787CIP2B 276	7756
626	1610	2594	3578	787CIP2B 277	7761
627	1611	2595	3579	787CIP2B 278	7761
628	1612	2596	3580	787CIP2B 279	7776
629	1613	2597	3581	787CIP2B 280	7783
630	1614	2598	3582	787CIP2B 281	7800
631	1615	2599	3583	787CIP2B 282	7800
632	1616	2600	3584	787CIP2B 283	7801
633	1617	2601	3585	787CIP2B 284	7811
634	1618	2602	3586	787CIP2B 285	7817
635	1619	2603	3587	787CIP2B 286	7821
636	1620	2604	3588	787CIP2B 287	7822
637	1621	2605	3589	787CIP2B 288	7841
638	1622	2606	3590	787CIP2B 289	7847
639	1623	2607	3591	787CIP2B 290	7880
640.	1624	2608	3592	787CIP2B 291	7910
641	1625	2609	3593	787CIP2B 293	7936
642	1626	2610	3594	787CIP2B 294	7945
643	1627	2611	3595	787CIP2B 295	7948
644	1628	2612	3596	787CIP2B 296	7963
645	1629	2613	3597	787CIP2B 297	7984
646	1630	2614	3598	787CIP2B 298	7985
647	1631	2615	3599	787CIP2B 299	8014
648	1632	2616	3600	787CIP2B 301	8029
649	1633	2617	3601	787CIP2B 302	8043
650	1634	2618	3602	787CIP2B_303	8164
651	1635	2619	3603	787CIP2B 304	8175
652	1636	2620	3604	787CIP2B 305	8250
653	1637	2621	3605	787CIP2B_306	8253
	1037			1	

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787CIP2B 363

787CIP2B 364

787CIP2B 365

787CIP2B 366

714	1698	2682	3666	787CIP2B 367	8646
715	1699	2683	3667	787CIP2B 368	8657
716	1700	2684	3668	787CIP2B 369	8661
717	1701	2685	3669	787CIP2B 370	8670
718	1702	2686	3670	787CIP2B 371	8692
719	1703	2687	3671	787CIP2B 372	8698
720	1704	2688	3672	787CIP2B 373	8762
721	1705	2689	3673	787CIP2B 374	8768
722	1706	2690	3674	787CIP2B 375	8768
723	1707	2691	3675	787CIP2B 376	8799
724	1708	2692	3676	787CIP2B 377	8806
725	1709	2693	3677	787CIP2B 378	8809
726	1710	2694	3678	787CIP2B 379	8814
727	1711	2695	3679	787CIP2B 380	8822
728	1712	2696	3680	787CIP2B 381	8833
729	1713	2697	3681	787CIP2B 382	8835
730	1714	2698	3682	787CIP2B 383	8877
731	1715	2699	3683	787CIP2B 384	8886
732	1716	2700	3684	787CIP2B 385	9003
733	1717	2701	3685	787CIP2B 386	9157
734	1718	2702	3686	787CIP2B 387	9175
735	1719	2703	3687	787CIP2B 388	9205
736	1720	2704	3688	787CIP2B 389	9260
737	1721	2705	3689	787CIP2B 390	9295
738	1722	2706	3690	787CIP2B 391	9307
739	1723	2707	3691	787CIP2B 392	9307
740	1724	2708	3692	787CIP2B 393	9312
741	1725	2709	3693	787CIP2B 394	9347
742	1726	2710	3694	787CIP2B_395	9370
743	1727	2711	3695	787CIP2B_396	9370
744	1728	2712	3696	787CIP2B_397	9382
745	1729	2713	3697	787CIP2B_398	9591
746	1730	2714	3698	787CIP2B_399	9650
747	1731	2715	3699	787CIP2B_400	9655
748	1732	2716	3700	787CIP2B_401	9663
749	1733	2717	3701	787CIP2B_402	9715
750	1734	2718	3702	787CIP2B_403	9755
751	1735	2719	3703	787CIP2B_404	9766
752	1736	2720	3704	787CIP2B_405	9771
753	1737	2721	3705	787CIP2B_406	9784
754	1738	2722	3706	787CIP2B_407	9925
755	1739	2723	3707	787CIP2B_408	9970
756	1740	2724	3708	787CIP2B_409	9997
757	1741	2725	3709	787CIP2B_410	10008
758	1742	2726	3710	787CIP2B_411	10010
759 760	1743	2727	3711	787CIP2B_412 787CIP2B_413	
	1744	2728	3712		10043
761	1745	2729	3713	787CIP2B_414	10093
762 763	1746	2730	3714	787CIP2B_415 787CIP2B_416	10172
764	1747	2731	3715	787CIP2B 416	10205
765	1748	2732	3716	787CIP2B_417 787CIP2B_418	10246
766	1749 1750		3717	787CIP2B_418 787CIP2B_419	10246
767	1751	2734	3718 3719	787CIP2B_419	886
768		2735		787CIP2C_1 787CIP2C_2	1028
769	1752	2736	3720	787CIP2C_2	1916
770	1753 1754		3721 3722	787CIP2C_3	2072
771	1754	2738	3722	787CIP2C_4	2424
772	1756	2740	3724	787CIP2C_5	2474
773	1757	2741	3725	787CIP2C_7	2474
	1131	12/71	13/23	70701120_7	1277

774	1758	2742	3726	787CIP2C 8	2887
775	1759	2743	3727	787CIP2C_8	3001
776	1760	2744	3728	787CIP2C_9	3182
777	1761	2745	3729	787CIP2C_10	3182
778	1762	2746	3730	787CIP2C_11	3182
779	1762	2747	3731	787CIP2C_12	3193
780	1764	2748	3732	787CIP2C_13	3196
781	1765	2749	3733	787CIP2C_14	3224
782	1766	2750	3734	787CIP2C_15	3225
783	1767	2751	3734	787CIP2C_16	3234
784	1768	2752	3736	787CIP2C_17	3241
785	1769	2753	3737	787CIP2C_18	3241
786	1770	2754	3738	787CIP2C_19	3243
787	1771	2755	3739	787CIP2C_20 787CIP2C_21	3259
			3740	787CIP2C_21	
788	1772	2756			3272
789	1773	2757	3741	787CIP2C_23	3278
790	1774	2758	3742	787CIP2C_24	3296
791	1775	2759	3743	787CIP2C_25	3327
792	1776	2760	3744	787CIP2C_26	3334
793	1777	2761	3745	787CIP2C_27	3339
794	1778	2762	3746	787CIP2C_28	3347
795	1779	2763	3747	787CIP2C_29	3387
796	1780	2764	3748	787CIP2C_30	3392
797	1781	2765	3749	787CIP2C_31	3411
798	1782	2766	3750	787CIP2C_32	3427
799	1783	2767	3751	787CIP2C_33	3432
800	1784	2768	3752	787CIP2C_34	3441
801	1785	2769	3753	787CIP2C_35	3479
802	1786	2770	3754	787CIP2C_36	3488
803	1787	2771	3755	787CIP2C_37	3488
804	1788	2772	3756	787CIP2C_38	3553
805	1789	2773	3757	787CIP2C_39	3560
806	1790	2774	3758	787CIP2C_40	3618
807	1791	2775	3759	787CIP2C_41	3642
808	1792	2776	3760	787CIP2C_42	3649
809	1793	2777	3761	787CIP2C_43	3676
810	1794	2778	3762	787CIP2C_44	3747
811	1795	2779	3763	787CIP2C_45	3917
812	1796	2780	3764	787CIP2C_46	4218
813	1797	2781	3765	787CIP2C_47	4219
814	1798	2782	3766	787CIP2C_48	4222
815	1799	2783	3767	787CIP2C_49	4222
816	1800	2784	3768	787CIP2C_50	4229
817	1801	2785	3769	787CIP2C_51	4230
818	1802	2786	3770	787CIP2C_52	4240
819	1803	2787	3771	787CIP2C_53	4241
820	1804	2788	3772	787CIP2C_54	4249
821	1805	2789	3773	787CIP2C_55	4252
822	1806	2790	3774	787CIP2C_56	4267
823	1807	2791	3775	787CIP2C_57	. 4272
824	1808	2792	3776	787CIP2C_58	4273
825	1809	2793	3777	787CIP2C_59	4275
826	1810	2794	3778	787CIP2C_60	4283
827	1811	2795	3779	787CIP2C_61	4290
828	1812	2796	3780	787CIP2C_62	4292
829	1813	2797	3781	787CIP2C_63	4305
830	1814	2798	3782	787CIP2C_64	4306
831	1815	2799	3783	787CIP2C_65	4308
832	1816	2800	3784	787CIP2C_66	4322
833	1817	2801	3785	787CIP2C_67	4351

		10000	10506	1 00=07000	1056	
834	1818	2802	3786	787CIP2C_68	4356 4399	
835	1819	2803	3787 3788	787CIP2C_69 787CIP2C_70	4400	
836 837	1820 1821	2805	3789	787CIP2C_70	4520	·
838	1822	2806	3790	787CIP2C 72	4598	
839	1823	2807	3791	787CIP2C_72	4599	
840	1824	2808	3792	787CIP2C_73	4600	
	1825	2809	3793	787CIP2C_74	4670	
841 842	1826	2810	3794	787CIP2C_73	4708	
843	1827	2811	3795	787CIP2C_76	4734	
844	1828	2812	3796	787CIP2C 78	4738	
845	1829	2813	3797	787CIP2C_78	4749	
846	1830	2814	3798	787CIP2C 80	4752	
847	1831	2815	3799	787CIP2C_80	4752	
848	1832	2816	3800	787CIP2C_81	4770	
849	1833	2817 ·		787CIP2C_82	4784	
850	1834	2818	3802	787CIP2C_83	4785	
851	1835	2819	3803	787CIP2C_85	4792	
		2820	3804	787CIP2C_85	4803	
852 853	1836 1837	2821	3805	787CIP2C_80	4811	
853 854	1837	2822	3806	787CIP2C_87	4817	
855	1839	2823	3807	787CIP2C 89	4818	
856	1840	2824	3808	787CIP2C_89	4820	
857	1841	2825	3809	787CIP2C_90	4831	
858	1842	2826	3810	787CIP2C_91	4841	
859	1843	2827	3811	787CIP2C_92	4869	
860	1844	2828	3812	787CIP2C_93	4876	
861	1845	2829	3813	787CIP2C 95	4902	
862	1846	2830	3814	787CIP2C 96	4910	
863	1847	2831	3815	787CIP2C 97	4931	
864	1848	2832	3816	787CIP2C 98	5303	
865	1849	2833	3817	787CIP2C 99	5317	
866	1850	2834	3818	787CIP2C 100	5322	
867	1851	2835	3819	787CIP2C 101	5330	
868	1852	2836	3820	787CIP2C 102	5333	
869	1853	2837	3821	787CIP2C_103	5333	
870	1854	2838	3822	787CIP2C 104	5356	
871	1855	2839	3823	787CIP2C 105	5363	
872	1856	2840	3824	787CIP2C 106	5364	
873	1857	2841	3825	787CIP2C 107	5379	
874	1858	2842	3826	787CIP2C 108	5386	
875	1859	2843	3827	787CIP2C 109	5397	
876	1860	2844	3828	787CIP2C 110	5401	
877	1861	2845	3829	787CIP2C 111	5419	
878	1862	2846	3830	787CIP2C 112	5420	
879	1863	2847	3831	787CIP2C 113	5452	
880	1864	2848	3832	787CIP2C 114	5467	
881	1865	2849	3833	787CIP2C 115	5482	
882	1866	2850	3834	787CIP2C 116	5483	
883	1867	2851	3835	787CIP2C 117	5492	
884	1868	2852	3836	787CIP2C 118	5499	
885	1869	2853	3837	787CIP2C 119	5525	
886	1870	2854	3838	787CIP2C 120	5538	
887	1871	2855	3839	787CIP2C 121	5539	
888	1872	2856	3840	787CIP2C 122	5558	-
889	1873	2857	3841	787CIP2C 123	5559	
890	1874	2858	3842	787CIP2C 124	5586	`
891	1875	2859	3843	787CIP2C 125	5619	
892	1876	2860	3844	787CIP2C 126	5628	
893	1877	2861	3845	787CIP2C 127	5640	

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TY O VALUE AZV

904	1970	2862	3846	787CIP2C 128	5640
894 895	1878 1879	2863	3847	787CIP2C_128	5827
896	1880	2864	3848	787CIP2C 130	6094
897	1881	2865	3849	787CIP2C 131	6195
898	1882	2866	3850	787CIP2C 132	6206
899	1883	2867	3851	787CIP2C 133	6355
	1884	2868	3852	787CIP2C 134	6362
900 901	1885	2869	3853	787CIP2C 135	6386
901	1886	2870	3854	787CIP2C 136	6431
902	1887	2871	3855	787CIP2C 137	6457
903	1888	2872	3856	787CIP2C 138	6480
904	1889	2873	3857	787CIP2C 139	6497
905	1890	2874	3858	787CIP2C 140	6532
907	1891	2875	3859	787CIP2C 141	6598
907	1892	2876	3860	787CIP2C_141	6644
		2877	3861	787CIP2C 143	6644
909	1893	2878	3862	787CIP2C 144	6645
910	1894		3863	787CIP2C_144	6645
911	1895	2879		787CIP2C_145	6761
912	1896	2880	3864	787CIP2C_146	6782
913	1897	2881	3865	787CIP2C_147	6981
914	1898	2882	3866 3867	787CIP2C_148 787CIP2C_149	6981
915	1899	2883		787CIP2C_149	7000
916	1900	2884	3868	787CIP2C_130	7029
917	1901	2885	3869	787CIP2C_131	7885
918	1902	2886	3870 3871	787CIP2C_132	8143
919	1903	2887	3872	787CIP2C_153	8143
920	1904		3873	787CIP2C 155	8234
921	1905	2889	3874	787CIP2C_155	8463
922	1906	2890	3875	787CIP2C_130	8467
923	1907	2891	3876	787CIP2C_157	8540
924	1908	2892 2893	3877	787CIP2C 159	8600
925	1909	2894	3878	787CIP2C 160	9656
926	1910	2895	3879	787CIP2C_100	9669
927	1911	2896	3880	787CIP2C 162	9695
928 929	1912 1913	2897	3881	787CIP2C_163	9744
	1913	2898	3882	787CIP2C 164	9849
930 931	1914	2899	3883	787CIP2D 1	4180
931	1915	2900	3884	787CIP2D 2	4181
	1917	2900	3885	787CIP2D_2	4314
933 934	1917	2901	3886	787CIP2D_3	4500
934	1918	2902	3887	787CIP2D_4	5651
936	1919	2903	3888	787CIP2D 6	5691
937	1920	2904	3889	787CIP2D_7	5881
937	1921	2905	3890	787CIP2D_7	5882
938	1922	2907	3891	787CIP2D_8	6209
939	1923	2907	3892	787CIP2D_9	6719
	1924	2908	3893	787CIP2D_10	8130
941			3894	787CIP2D_11	8863
942	1926 1927	2910 2911	3894	787CIP2D_12 787CIP2D 13	8902
943	1927	2911	3896	787CIP2D_13	9162
944			3896	787CIP2D_14 787CIP2D_15	9197
945	1929	2913		787CIP2D_13	9215
946	1930	2914	3898	787CIP2D_16	9232
947	1931	2915	3899	787CIP2D_17 787CIP2D_18	9232
948	1932	2916	3900	787CIP2D_18 787CIP2D_19	9369
949	1933	2917	3901	787CIP2D_19 787CIP2D 20	
950	1934	2918	3902	L	9371
951	1935	2919	3903	787CIP2D_21	9516
952	1936	2920	3904	787CIP2D_22	9601
953	1937	2921	3905	787CIP2D_23	9731

TI W WAIM LASH

954	1938	2922	3906	787CIP2D_24	9733
955	1939	2923	3907	787CIP2D_25	9769
956	1940	2924	3908	787CIP2D_26	9804
957	1941	2925	3909	787CIP2D_27	9816
958	1942	2926	3910	787CIP2D_28	9844
959	1943	2927	3911	787CIP2D_29	9924
960	1944	2928	3912	787CIP2D_30	9936
961	1945	2929	3913	787CIP2D_31	10163
962	1946	2930	3914	787CIP2D_32	10165
963	1947	2931	3915	787CIP2D_33	10165
964	1948	2932	3916	787CIP2D_34	10244
965	1949	2933	3917	787CIP2D_35	10278
966	1950	2934	3918	787CIP2E_1	4251
967	1951	2935	3919	787CIP2E_2	5310
968	1952	2936	3920	787CIP2E_3	5697
969	1953	2937	3921	787CIP2E_4	5731
970	1954	2938	3922	787CIP2E_5	5733
971	1955	2939	3923	787CIP2E_6	5734
972	1956	2940	3924	787CIP2E_7	5740
973	1957	2941	3925	787CIP2E_8	7657
974	1958	2942	3926	787CIP2E_9	9572
975	1959	2943	3927	787CIP2F_1	1363
976	1960	2944	3928	787CIP2F_2	4303
977	1961	2945	3929	787CIP2F_3	5760
978	1962	2946	3930	787CIP2F_4	5766
979	1963	2947	3931	787CIP2F_5	5767
980	1964	2948	3932	787CIP2F_6	5767
981	1965	2949	3933	787CIP2F_7	5770
982	1966	2950	3934	787CIP2F_8	6855
983	1967	2951	3935	787CIP2F_9	10026
984	1968	2952	3936	787CIP2F_10	10227

TABLE 6

SEQ ID NO:	Method	Predicted beginning nucleotide location corresponding to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, I=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion
2953	A	3	324	ISEHRIEASGNYLAQRLTSSFLRGLSSWKSNPLML CGWTILLTLTMVQGEP*GP\KGIPG\FHTNSSYPH WGTVAKPPAGD*DLLPAPGQEGTPLFTR*SLCTY CPID
2954	A	18	467	REELGKDLFDCTLYVLLKYDDFNADKHLALEEF YRAFQVIQLSLPEDQKLSITAATVGQSAVLSCAIQ GTLRPPIIWKRNNIILNNLDLEDINDFGDDGSLYIT KVTTTHVGNYTCYADGYEQVYQTHIFQVNVPPV IRVYPESQARRAG
2955	A	3	23	FYSAFLVADKGIVTSKHNNDTQHIWESDSNEFSV IADPRGNTLGRGTTIT*VSIPPSL
2956	A	1	493	RTKTDVYILNLAVADLLLLFTLPFWAVNAVHGW VLGKIMCKITSALYTLNFVSGMQFLACISIDRYV AVTKVPSQSGVGKPCWIICFCVWMAAILLSIPQL VFYTVNDNARCIPIFPRYLGTSMKALIQMLEICIG FVVPFLIMGVCYFITARTLMKMPNIKIS
2957	A	703	302	EETGVREKRRERMKEKMWQNVLCCTLQTAVIL KLFQNKVLNILKNFFLSPLDTRKNKVFKKWAGG PGAVAHACNPSTLGGRGGRITKSGDRDHPGQHG

	T 5 2 - 5 -		T 100 101	
SEQ ID NO:	Method	Predicted beginning nucleotide location corresponding to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion
				ETRSLPACWAQWKSLALPVSRAPGRQGSLVVFP LP
2958	A	575	1054	CTKCKADCDTCFNKNFCTKCKSGFYLHLGKCLD NCPEGLEANNHTMECVSIVHCEVSEWNPWSPCT KKGKTCGFKRGTETRVREIIQHPSAKGNLCPPTN ETRKCTVQRKKCQKGERGKKGRERKRKKPNKG ESKEAIPDSKSLESSKEIPEQRENKQQQ
2959	A	1	426	LSMLSTISTEHRLSVLWPIWYCCHCPTHLSAVMC VLLWALSLLQSILEWMFCSFLFSDVDSDNWCQIL DFLTAVWLIFLI\LVLCGFTLVLLVRIICGSQKMPL TRLYVTILLTGLVFLFCSLPLSIQ*FLLYWIEKDLD DL
2960	A .	1194	852	EKRKTSYSQCLNSKQRNVSMRPSIWIHVHLKPPC RLVELLPFSSALQGLSHLSLGTTLP/V*GHLRFRL RNLPQSLRTVILPERNEEQNLQELSHNADKYQM GDCCKEEIDDSIFY
2961	A	274	2250	EKGKVKDAGAEQWISLSLSCKGSWETQFSNHLN SLTPPTSVRRMPLITTVTLLKMVARHHMKLLCSK AFSTQLQQKIFLHSQMGIHHQSVCMKLKPNTSHII SILMGQPMALVQLETLAPLTIIIQKFQTQDHMKF WKNLPLHSHHLTPSVPQTVIPKKTGSPEIKLKITK TIQNGRELFESSLCGDLLNEVQASE\Q*NQSIESRK EKRKKSNKHDSSRSEERKSHKIPKLEPEEQNRPN ERVDTVSEKPREEPVLKEGSPSSANTIFCSNNGSV HWFKFQVGDLVWSKVGTYPWWPCMVSSDPQL EVHTKINTRGAREYHVQFFSNQPERAWVHEKRV REYKGHKQYEELLAEATKQASNHSEKQKIRKPR PQRERAQWDIGIAHAEKALKMTREERIEQYTFIYI DKQPEEALSQAKKSVASKTEVKKTRPRSVLNT QPEQTNAGEVASSLSSTEIRRHSQRRHTSAEEEEP PPVKIAWKTAAARKSLPASITMHKGSLDLQKCN MSPVVKIEQVFALQNATGDGKFIDQFVYSTKGIG NKTEISVRGQDRLIISTPNQRNEKPTQSVSSPEATS GSTGSVEKKQQRRSIRTRSESEKSTEVVPKKKIK KEQVETVPQATVKTGLQKGSADRGVQGSVRFSD SSVSAAIEETVD
2962	A	2408	836	SASPPPPPPPPSRFPFSGAPGARDRSGPLGSEPQR NPGARPRTLEATVTPPGSVGAMSSSGLNSEKVA ALIQKLNSDPQFVLAQNVGTTHDLLDICLKRATV QRAQHVFQHAVPQEGKPITNQKSSGRCWIFSCLN VMRLPFMKKLNIEEFEFSQSYLFFWDKVERCYFF LSAFVDTAQRKEPEDGRLVQFLLMNPANDGGQ WDMLVNIVEKYGVIPKKCFPESYTTEATRRMND ILNHKMREFCIRLRNLVHSGATKGEISATQDVM MEEIFRVVCICLGNPPETFTWEYRDKDKNNKKIG P\ITPLEFNR/EQHVKPLFNMEDKICLVNDPRPQH KYNKLYTV\EYL\SNMVWRGEKLFYNNQPIDFLK KMVAASIKDG\EAVWFGCDVGKHF\NSKLG\LSD MNLYDHELVFGVSLKNMNKAER\LTFGES\LMT HTMTFTAV/SQSRDDSGMVLFTKW\RVGEFQWG EDHGH\KGYLCMTD*VGSLEYVYEVV/VWDRKH VP\EEVLAVLGAGNPFVLPAWDPMGALAE
2963	A	90	543	RHYDSAGKITLKIAKNYLEQRAVGGASPRLAQS VLTCSREPILENSLTSLIEYLHNALEHDMRLRFNN DRMKTTIKETST*LSNSYLVFPLM*SLTYLMKMS

Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methlonine, Predicted end SEQ ID Method Predicted beginning nucleotide location NO:

No.		nucleotide location corresponding to first amino acid residue of peptide sequence	location corresponding to last amino acid residue of peptide sequence	I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion
				FERCTARNKMFVNSPFTKVDNYCT\SS\WKKFYL KCYFSLNTIKKEKKMT
2964	A		2454	FDTYRGLPSISNGNYSQLQFQAREYSGAPYSQRIS AITTVSVAWKVLSGKIGEGAEGNCKCVISEGAW AVCPTQPCGKAKPDKHLKDLLSKLLNSGYFESIP VPKNAKEKEVPLEEEMLIQSEKKTQLSKTESVKE SESLMEFAQPEIQPQEFLNRRYMTEVDYSNKQGE EQPWEADYARKPNLPKRWDMLTEPDGQEKKQE SFKSWEASGKHQEVSKPAVSLEQRKQDTSKLRS TLPEEQKKQEISKSKPSPSQWKQDTPKSKAGYVQ EEHKKQETPKLWPVQLQKEQDPKKQTPKSWENN VESQKHSLTSQSQISPKSWGVATASLIPNDQLLPR KLNTEPKDVP/IACASA*GFLPLQPPFRRI/HVLRK EKLQDLMTQIQGTCNFMQESVLDFDKPSSAIPTS QPPSATPG*PRRHLKEQNLS\VKVIFFQGAVT\VF NVNAPLPPRKEQEIKESPYSPGYNQSFTTASTQTP PQCQLPSIHVEQTVHSQETANYHPDGTIQVSNGS LAFYPAQTNVFPRPTQPFVNSRGSVRGCTRGGRL ITNSYRSPGGYKGFDTYRGLPSISNGNYSQLQFQ AREYSGAPYSQRDNFQQCYKRGGTSGGPRANSR AGWSDSSQVSSPERDNETFNSGDSGQGDSRSMT PVDVPVTNPAATILPVHVYPLPQQMRVAFSAAR TSNLAPGTLDQPIVFDLLLNNLGETFDLQLGRFN CPVNGTYVFIFHMLKLAVNVPLYVNLMKNEEVL VSAYANDGAPDHETASNHAILQLFQGDQIWLRL HRGAIYGSSW
2965		3	2454	FDTYRGLPSISNGNYSQLQFQAREYSGAPYSQRIS AITTVSVAWKVLSGKIGEGAEGNCKCVISEGAW AVCPTQPCGKAKPDKHLKDLLSKLLNSGYFESIP- VPKNAKEKEVPLEEEMLIQSEKKTQLSKTESVKE SESLMEFAQPEIQPQEFLNRRYMTEVDYSNKQGE EQPWEADYARKPNLPKRWDMLTEPDGQEKKQE SFKSWEASGKHQEVSKPAVSLEQRKQDTSKLRS TLPEEQKKQEISKSKPSPSQWKQDTPKSKAGYVQ EEHKKQETPKLWPVQLQKEQDPKKQTPKSWENN VESQKHSLTSQSQISPKSWGVATASLIPNDQLLPR KLNTEPKDVP/IACASA*GFLPLQPPFRRI/HVLRK EKLQDLMTQIQGTCNFMQESVLDFDKPSSAIPTS QPPSATPG*PRRHLKEQNLS\VKVIFFQGAVT\VF NVNAPLPPRKEQEIKESPYSPGYNQSFTTASTQTP PQCQLPSIHVEQTVHSQETANYHPDGTIQVSNGS LAFYPAQTNVFPRPTQPFVNSRGSVRGCTRGGRL ITNSYRSPGGYKGFDTYRGLPSISNGNYSQLQFQ AREYSGAPYSQRDNFQQCYKRGGTSGGPRANSR AGWSDSSQVSSPERDNETFNSGDSGQGDSRSMT PVDVPVTNPAATILPVHVYPLPQQMRVAFSAAR TSNLAPGTLDQPIVFDLLLNNLGETFDLQLGRFN CPVNGTYVFIFHMLKLAVNVPLYVNLMKNEEVL VSAYANDGAPDHETASNHAILQLFQGDQIWLRL HRGAIYGSSW
2966	A	1693	227	DYVLTAELHRQRSPGVSFGLSVFNLMNAIMGSGI LGLAYVMANTGVFGFSFLLLTVALLASYSVHLL LSMCIQTAYLGP*TNYFMVLPAH*LTCLPLIEFLQ

				2 (2) (((((((((((((((((((((((((((((((((
SEQ ID NO:	Method	Predicted beginning nucleotide location corresponding to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \text{possible nucleotide insertion}
	•	·		SL*NSL*AVTSYEDLGLFAFGLPGKLVVAGTIIIQ NIGAMSSYLLIIKTELPAAIAEFLTGDYSRYWYLD GQTLLIIICVGIVFPLALLPKIGFLGYTSSLSFFFM MFFALVVIIKKWSIPCPLTLNYVEKGFQISNVTDD CKPKLFHFSKESAYALPTMAFSFLCHTSILPIYCE LQSPSKKRMQNVTNTAIALSFLIYFISALFGYLTF YD/GTTKAQRGEVTCHRIKDKVESELLKG***IP* SHDVVVMT\VKLCILFAVLL\TVPLIHFPARKAVT MMFFSNFPFSWIRHFLITLALNIIIVLLAIYVPDIRN VFGVVGASTSTCLIFIFPGLFYLKLSREDFLSWKK LGVGCFC/LLSFKTSILRNSLSVYIILPASRKSIYFK I
2967		3	3222	SGIVVRALWREKKPGGGRRVKRRNPGRQAVGH TEEDPPRVGTPWKEHTGPGPQEGSTMEAAHAKT TEECLAYFGVSETTGLTPDQVKRNLEKYGLNELP AEEGKTLWELVIEQFEDLLVRILLAACISFVLA WFEEGEETITAFVEPFVILLILIANAIVGVWQERN AENAIEALKEYEPEMGKVYRADRKSVQRIKARD IVPGDIVEVAVGDKVPADIRILAIKSTTLRVDQSIL TGEYVSVIKHTEPVPDPRAVNQDKKNMLFSGTNI AAGKALGIVATTGVGTEIGKIRDQMAATEQDKT PLQQKLDEFGEQLSKVISLICVAVWLINIGHFNDP VHGGSWFRGAIYYFKIAVALAVAAIPEGLPAVIT TCLALGTRRMAKKNAIVRSLPSVETLGCTSVICS DKTGTLTINQMSVCKMFIIDKVDGDICLLNEFSIT GSTYAPEGEVLKNDKPVRPGQYDGLVELATICA LCNDSSLDFNEAKGVYEKVGEATETALTTLVEK MNVFNTDVRSLSKVERANACNSVIRQLMKKEFT LEFSRDRKSMSVYCSPAKSSRAAVGNKMFVKGA. PEGVIDRCNYVRVGTTRVPLTGPVKEKIMAVIKE. WGTGRDTLRCLALATRDTPPKREEMVLDDSARF LEYETDLTFVGVVGMLDPPRKEVTGSIQLCRDA GIRVIMITGDNKGTAIAICRRIGIFGENEEVADRA YVTGREFDDLVPLAEQNEACRRACCFARVEPSHK SKIVEYLQSYDEITAMTGDGVNDAPALKKAEIGI AMGSGTAVAKTASEMVLADDNFSTIVAAVEEGR AIYNNMKQFIRYLISSNVGEVVCIFLTAALGLPEA LIPVQLLWVNLVTDGLPATALGFNPPDLDIMDRP PRSPKEPLISGWLFFRYMAIGGYVGAATVGAAA WWFLYAEDGPHVNYSQLTHFMQCTEDNTHFEGI DCEVFEAPEPMTMALSVLVTIEMCNALNSLSEN QSLLRMPPWVNIWLLGSICLSMSLHFLILYVDPLP MIFKLRALDLTQWLMVLKISLPVIGLDEILKFVA RNYLEG*LFPLLHL*ARVTDPEDERRK
2968	A	3	2414	GARSCSRLGRCTFPLWKGREMEVRKLSISWQFLI VLVLILQILSALDFDPYRVLGVSRTASQADIKKA YKKLAREWHPDKNKDPGAEDKFIQISKAYEILSN EEKRSNYDQYGDAGENQGYQKQQQREYRFRH FHENFYFDESFFHFPFNSERRDSIDEKYLLHFSHY VNEVAPDSFKKPYLIKITSDWCFSCIHIEPVWKEV IQELEELGVGIGVVHAGYERRLAHHLGAHSTPSI LGIINGKISFFHNAVVRENLRQFVESLLPGNLVEK VTNKNYVRFLSGWQQENKPHVLLFDQTPIVPLL YKLTAFAYKDYLSFGYVYVGLRGTEEMTRRYNI NIYAPTLLVFKEHINRPADVIQARGMKKQIIDDFI

SEQ ID NO:	Method	Predicted beginning nucleotide location corresponding to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \(\=\text{possible nucleotide insertion} \)
			-	TRNKYLLAARLTSQKLFHELCPVKRSHRQRKYC VVLLTAETTKLSKPFEAFLSFALANTQDTVRFVH VYSNRQQEFADTLLPDSEAFQGKSAVSILERRNT AGRVVYKTLEDPWIGSESDKFILLGYLDQLRKDP ALLSSEAVLPDLTDELAPVFLLRWFYSASDYISD CWDSIFHNNW\REMMPLLSLIFSALFILFGTVIVQ AFSDSNDERESSPPEKEEAQEKTGKTEPSFTKENS SKIPKKGFVEVTELTDVTYTSNLVRLRPGHMNV VLILSNSTKTSLLQKFALEVYTFTGSSCLHFSFLSL DKHREWLEYLLEFAQDAAPIPNQYDKHFMERDY TGYVLALNGHKKYFCLFKPQKTVEEGGKP*GSC SDVDSSLYLGESRGKPSCGLGSRPIKGKLSKLSL WMERLLEGSLQRFYIPSWPELD
2969	A	48	1117	KGLSPDQVLSAFAPLDCEMWLKVFTTFLSFATG ACSGLKVTVPSHTVHGVRGQALYLPVHYGFHTP ASDIQIIWLFERPHTMPKYLLGSVNKSVVPD/YGI P/YTSSP*CHPMASLLINPLQFPDEGNYIVKVNIQG NGTLSASQKIQVTVDDPVTKPVVQIHPPSGAVEY VGNMTLTCHVEGGTRLAYQWLKNGRPVHTSST YSFSPQNNTLHIAPVTKEDIGNYSCLVRNPVSEM ESDIIMPIIYYGPYGLQVNSDKGLKVGEVFTVDL GEAILFDCSADSHPPNTYSWIRRTDNTTYIIKHGP RLEVASEKVAQKTMDYVCCAYNNITGRQDETHF TVIITSVGMCDIQGRDPNKT
2970	A	68	936	HSALLTHSSFCVFTLCQDFFTYSSMSEEVTYADL QFQNSSEMEKIPEIGKFGEKAPPAPSHVWRPAAL FLTLLCLLLLIGLGVLASMFHVTLKIEMKKMNKL QNISEELQRNISLQLMSNMNISNKIRNLSTTLQTI ATKLCRELYSKEQEHKCKPCPRRWIWHKDSCYF LSDDVQTWQESKMACAAQNASLLKINNKNALE FIKSQSRSYDYWLGLSPEEDS/YSWYESG*YNQ\P SAWVIRNAPDLNNMYCGYINRLYVQYYHCTYK QRMICEKMANPVQLGSTYFREA
2971	A	912	2287	VPNYLPSVSSAIGGEVPQRYVWRFCIGLHSAPRF LVAFAYWNHYLSCTSPCSCYRPLCRLNFGLNVV ENLALLVLTYVSSSEDF/TWVPG*GRSGEVFPEGT GLPLPHSDLPTSWCGHSLQCGSQSSFPPAIHENAF IVFIASSLGHMLLTCILWRLTKKHTVSQE\DGLSL AGAPRQPRRKSRTSVLRIRVMVRWELSSNGNPG RGVLGLGLGLGNKLRVVGQNLGL*HCVWVVWE TGE*KRWRLQMGIE*GVASRRQ*VRNSVRGLVC HNSSAPPMYMGFFSPTVFGGGVGG*LHVTFILHP PEVEAAGIPLLLGPSLPQRQGREHIVVILAAPACA PFHDR*WEPREIRPSP*ELGLRGEPTLSYPASCRVI RQPIP*DRKSYSWKQRLFIINFISFFSALAVYFRHN MYCEAGVYTIFAILEYTVVLTNMAFHMTAWWD FGNKELLITSQPEEKRF
2972	A	1734	246	GGILSGRDGRTALPRPREPAERTAGLRRDMRPQE LPRLAFPLLLLLLLLPPPPCPAHSATRFDPTWES LDARQLPAWFDQAKFGIFIHWGVFSVPSFGSEWF WWYWQKEKIPKYVEFMKDNYPPSFKYEDFGPL FTAKFFNANQ\WADIFQASGAKYIVLTSKHHEGF TLWG\SEYSWNWNAIDEGPKRDIVKELEVAIRNR TDLRFGLYYSLFEWFHPLFLEDESSSFHKRQFPVS KTLPELYELVNNYQPEVLWSDGDGGAPDQYWN

SEQ ID NO:	Method	Predicted beginning nucleotide location corresponding to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion
				STGFLAWLYNESPVRGTVVTNDRWGAGSICKHG GFYTCSDRYNPGHLLPHKWENCMTIDKLSWGY RREAGISDYLTIEELVKQLVETVSCGGNLLMNIG PTLDGTISVVFEERLRQMGSWLKVNGEAIYETHT WRSQNDTVTPDVWYTSKPKEKLVYAIFLKWPTS GQLFLGHPKAILGATEVKLLGHGQPLNWISLEQN GIMVELPQLTIHQMPCKWGWALALTNVI
2973	A		1133	SVPRAGGDMETGAAELYDQALLGILQHVGNVQ DFLRVLFGFLYRKTDFYRLLRHPSDRMGFPPGAA QALVLQVFKTFDHMARQDDEKRRQELEEKIRRK EEEEAKTVSAAAAEKEPVPVPVQEIEIDSTTELDG HQEVEKVQPPGPVKEMAHGSQEAEAPGAVAGA AEVPR\EPPILPRIQEQFQKNPDSYNGAVRENYTW SQDYTDLEVRVPVPKHVVKGKQVSVALSSSSIRV AMLEENGERVLMEGKLTHKINTESSLWSLEPGK CVLVNLSKVGEYWWNAILEGEEPIDIDKINKERS MATVDEEEQAVLDRLTFDYHQKLQGKPQSHEL KVHEMLKKGWDAEGSPFRGQRFDPAMFNISPGA VQF
2974	A	271	1854	MQFGRAHGDCVSGAQLCGCPSMDDYMVLRMIG EGSFGRALLVQHESSNQMFAMKEIRLPKSFSNTQ NSRKEAVLLAKMKHPNIVAFKESFEAEGHLYIV MEYCDGGDLMQKIKQQKGKLFPEDMILNWFTQ MCLGVNHIHKKRVLHRDIKSKNIFLTQNGKGKL GDFGSARLLSNPMAFACTYVGTPYYVPPEIWEN LPYNNKSDIWSLGCILYELCTLKHPFQANSWKNL ILKVCQGCISPLPSHYSYELQFLVKQMFKRNPSH RPSATTLLSRGIVARLVQKCLPPEIIMEYGEEVLE EIKNSKHNTPRKKTNPSRIRIALGNEASTVQEEEQ DRKGSHTDLESINENLVESALRRVNREEKGNKSV. HLRKASSPNLHRRQWEKNVPNTALTALENASILT SSLTAEDDRGGSVIKYSKNTTRKQWLKETPDTLL NILKNADLSLAFQTYTIYRPGS\EGFLKGPLSEETE ASDSVDGGHDSVILDPERLEPGLDEEDTDFEEED DNPDWVSELKKRAGWQGLCDR
2975	A		2833	PPGEPGAGRGALSPCGPLSGPPPLPGREAGGTCG QPVNPVFDLSRRNPQEDFELIQRIGSGTYGDVYK ARNVNTGELAAIKVIKLEPGEDFAVVQQEIIMMK D\CKHP\DIVAYF\GSYL\RRDKLWI\CMEF\CGSGS \LQDIYHVTGPLSELQIAYVSRETLQGLYYLHSKG KMHRDIKGANILLTDNGHVKLADFGVSAQITATI AKRKSFIGTPYWMAPEVAAVERKGGYNQLCDL WAVGITAIELAELQPPMFDLHPMRALFLMTKSNF QPPKLKDKMKWSNSFHHFVKMALTKNPKKRPT AEKLLQHPFVTQHLTRSLAIELLDKVNNPDHSTY HDFDDDDPEPLVAVPHRIHSTSRNVREEKTRSEIT FGQVKFDPPLRKETEPHHELPDSDGFLDSSEEIYY TARSNLDLQLEYGQGHQG\GYFLGANKSLLKSV EEELHQRGHVAHLEDDEGDDDESKHSTLKAKIP PPLPPKPKSIFIPQEMHSTEDENQGTIKRCPMSGSP \AKPSQVPPRPPPPRLPPHKPVALGNGMSSFQLNG ERDGSLCQQQNEHRGENLSRKEKKDVPKPISNG LPPTPKVHMGACFSKVFNGCPLKIHCASSWINPD TRDQYLIFGAEEGIYTLNLNELHETSMEQLFPRR CTWLYVMNNCLLSISGKASQLYSHNLPGLFDYA

SEO ID	Method	Predicted	Predicted end	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid,
NO:	Method	beginning	nucleotide	E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine,
}		nucleotide	location	I=Isoleucine, K=Lysine, L=Leucine, M=Methionine,
	1	location	corresponding to last amino	N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine,
1		corresponding to first amino	acid residue of	X=Unknown, *=Stop codon, /=possible nucleotide deletion,
ĺ	ľ	acid residue of	peptide	\=possible nucleotide insertion
		peptide sequence	sequence	•
		bequee		RQMQKLPVAIPAHKLPDRILPRKFSVSAKIPETK
				WCQKCCVVRNPYTGHKYLCGALQTSIVLLEWV
				EPMQKFMLIKHIDFPIPCPLKMFEMLVVPEQEYP
			!	LVCVGVSRGRDFNQVVRFETVNPNSTSSWFTES
				DTPQTNVTHVTQLERDTILVCLDCCIKIVNLQGR
				LKSSRKLSSELTFDFRIESIVCLQDSVLAFWKHG MQGRSFRSNEVTQEISDSTRIFRLLGSDRVVVLES
				RPTDNPTANSNLYILAGHENSY
2076		122	2833	PPGEPGAGRGALSPCGPLSGPPPLPGREAGGTCG
2976	A	32	2033	QPVNPVFDLSRRNPQEDFELIQRIGSGTYGDVYK
				ARNVNTGELAAIKVIKLEPGEDFAVVQQEIIMMK
1				D\CKHP\DIVAYF\GSYL\RRDKLWI\CMEF\CGSGS
				LQDIYHVTGPLSELQIAYVSRETLQGLYYLHSKG
				KMHRDIKGANILLTDNGHVKLADFGVSAQITATI
			·	AKRKSFIGTPYWMAPEVAAVERKGGYNQLCDL
				WAVGITAIELAELQPPMFDLHPMRALFLMTKSNF
į				QPPKLKDKMKWSNSFHHFVKMALTKNPKKRPT
j			,	AEKLLQHPFVTQHLTRSLAIELLDKVNNPDHSTY
				HDFDDDDPEPLVAVPHRIHSTSRNVREEKTRSEIT
				FGQVKFDPPLRKETEPHHELPDSDGFLDSSEEIYY
{	ł			TARSNLDLQLEYGQGHQG\GYFLGANKSLLKSV
1				EEELHQRGHVAHLEDDEGDDDESKHSTLKAKIP
		ļ	ļ	PPLPPKPKSIFIPQEMHSTEDENQGTIKRCPMSGSP
	Ì			\AKPSQVPPRPPPPRLPPHKPVALGNGMSSFQLNG
				ERDGSLCQQNEHRGENLSRKEKKDVPKPISNG
				LPPTPKVHMGACFSKVFNGCPLKIHCASSWINPD
				TRDQYLIFGAEEGIYTLNLNELHETSMEQLFPRR
				CTWLYVMNNCLLSISGKASQLYSHNLPGLFDYA RQMQKLPVAIPAHKLPDRILPRKFSVSAKIPETK
İ				WCQKCCVVRNPYTGHKYLCGALQTSIVLLEWV
				EPMQKFMLIKHIDFPIPCPLKMFEMLVVPEQEYP
			[LVCVGVSRGRDFNQVVRFETVNPNSTSSWFTES
				DTPQTNVTHVTQLERDTILVCLDCCIKIVNLQGR
1				LKSSRKLSSELTFDFRIESIVCLQDSVLAFWKHG
			,	MQGRSFRSNEVTQEISDSTRIFRLLGSDRVVVLES
				RPTDNPTANSNLYILAGHENSY
2977	A	174	1543	YSLRKGITFKLAGAMVHIKKGELTQEEKELLEVI
		}		GKGTVQEAGTLLSSKNVRVNCLDENGMTPLMH
				AAYKGKLDMCKLLLRHGADVNCHQHEHGYTA
1		1	(LMFAALSGNKDITWVMLEAGAETDVVNSVGRT
				AAQMAAFVGQHDCVTIINNFFPRERLDYYTKPQ
		}		GLDKEPKLPPKLAGPLHKIITTTNLHPVKIVMLV
				NENPLLTEEAALNKCYRVMDLICEKCMKQRDM
			ļ	NEVLAMKMHYISCIFQKCINFLKDGENKLDTLIK
		1	ĺ	SLLKG\RASDGFPVYPEKILRESIRK\FPYCEATLL
				QQLVRSIAPVEIGSDPTAFSVLTQAITGQVGFVDV
				EFCTTCGEKGASKRCSVCKMVIYCDQTCQKTHW FTHKKICKNLKDIYEKQQLEAAKEKRQEENHGK
		1	ļ	LDVNSNCVNEEQPEAEVGISQKDSNPEDSGEGK
			1 ·	KESLESEAELEGLQDAPAGPQVSEE
2079	 	3	5177	SDDLRTGLFQDVQDAESLKLPGVYEVLFYNETE
2978	A	13	31//	SDDLRIGLIQUVQDAESLALFGVIEVLFINETE DCPGMMLWRYPEPRGLTLVRITPVPFNTTEDPDI
				STADLGDVLQDPCSLEYWDELQKVFVAFREFNL
		1		SESKVCELQLPDINLVNDQKKLVSSDLWRIVLNS
				SONGADDOSSASESGSOSTCDPLVTPTALAACTR
L	J	<u> </u>	L	- Alian Annia Anni

SEQ ID	Method	Predicted	Predicted end nucleotide	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine,
NO:		beginning nucleotide	location	I=Isoleucine, K=Lysine, L=Leucine, M=Methionine,
	}	location	corresponding	N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine,
	ł	corresponding	to last amino	T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine,
	1	to first amino	acid residue of	X=Unknown, *=Stop codon, /=possible nucleotide deletion,
		acid residue of	peptide	\=possible nucleotide insertion
		peptide	sequence	
		sequence		VDSCFTPWFVPSLCVSFQFAHLEFHLCHHLDQLG
	1		1	TAAPQYLQPFVSDRNMPSELEYMIVSFREPHMYL
				ROWNINGSVCQEIQFLAQADCKLLECRNVTMQS
			}	VVKPFSIFGQMAVSSDVVEKLLDCTVIVDSVFVN
ļ				LGQHVVHSLNTAIQAWQQNKCPEVEELVFSHFV
			ŧ I	ICNDTQETLRFGQVDTDENILLASLHSHQYSWRS
	İ			HKSPQLLHICIEGWGNWRWSEPFSVDHAGTFIRT
				IQYRGRTASLIIKVQQLNGVQKQIIICGRQIICSYL
				SQSIELKVVQHYIGQDGQAVVREHFDCLTAKQK
ŀ]			LPSYILENNELTELCVKAKGDEDWSRDVCLESK
				APEYSIVIQVPSSNSSIIYVWCTVLTLEPNSQVQQ
		į		RMIVFSPLFIMRSHLPDPIIIHLEKRSLGLSETQIIP
		Ì		GKGQEKPLQNIEPDLVHHLTFQAREEYDPSDCA
				VPISTSLIKQIATKVHPGGTVNQILDEFYGPEKSL
	}			QPIWPYNKKDSDRNEQLSQWDSPMRVKLSIWKP
į	1			YVRTLLIELLPWALLINESKWDLWLFEGEKIVLQ
	1			VPAGKIIIPPNFQEAFQIGIYWANTNTVHKSVAIK
				LVHNLTSPKWKDGGNGEVVTLDEEAFVDTEIRL
				GAFPGHQKLCQFCISSMVQQGIQIIQIEDKTTIINN
ŀ				TPYQIFYKPQLSVCNPHSGKEYFRVPDSATFSICP
ļ				GGEQPAMKSSSLPCWDLMPDISQSVLDASLLQK'
				QIMLGFSPAPGADSSQCWSLPAIVRPEFPRQSVA
]			VPLGNFRENGFCTRAIVLTYQEHLGVTYLTLSED
•				PSPRVIIHNRCPVKMLIKENIKDIPKFEVYCKKIPS
1.				ECSIHHELYHQISSYPDCKTKDLLPSLLLRVEPLD
		,		
				EVTTEWSDAIDINSQGTQVVFLTGFGYVYVDVV
				HQCGTVFITVAPEGKAGPILTNTNRAPEKIVTF/K
		:	ŀ	MFITQLSLAVFDDLTHHKASAELLRLTLDNIFLC
				VAPGAGPLPGEEPVAALFELYCVEICCGDLQLDN
ļ				QLYNKSNFHFAVLVCQGEKAEPIQCSKMQSLLIS
ļ				NKELEEYKEKCFIKLCITLNEGKSILCDINEFSFEL
				KPARLYVEDTFVYYIKTLFDTYLPNSRLAGHSTH
'				LSGGKQVLPMQVTQHARALVNPVKLRKLVIQPV
ļ	1			NLLVSIHASLKLYIASDHTPLSFSVFERGPIFTTAR
	1			QLVHALAMHYAAGALFRAGWVVGSLDILGSPA
				SLVRSIGNGVADFFRLPYEGLTRGPGAFVSGVSR
1				GTTSFVKHISKGTLTSITNLATSLARNMDRLSLDE
				EHYNRQEEWRRQLPESLGEGLRQGLSRLGISLLG
				AIAGIVDQPMQNFQKTSEAQASAGHKAKGVISG
	1			VGKGIMGVFTKPIGGAAELVSQTGYGILHGAGLS
	1			QLPKQRHQPSD\VHADQAPNSHVKYVWKMLQS
			1.	LGRPEVHMALDVVLVRGSGQEHEGCLLLTSEVL
1	1		1	FVVSVSEDTQQQAFPVTEIDCAQDSKQNNLLTV
]		QLKQPRVACDVEVDGVRERLSEQQYNRLVDYIT
			·	KTSCHLAPSCSSMQIPCPVVAAEPPPSTVKTYHY
				LVDPHFAQVFLSKFTMVKNKALRKGFP
2979	A	255	2673	AWLFPASVLCPRCLTGSAVGSAEWKSLVVLFPFS
2713	/ h	233	20,3	SRPTLGHLDSKPSSKSNMIRGRNSATSADEQPHIG
1				NYRLLKTIGKGNFAKVKLARHILTGKEVAVKIID
		{		KTQLNSSSLQKLFREVRIMKVLNHPNIVKLFEVIE
		}		TEKTLYLVMEYASGGEVFDYLVAHGRMKEKEA
				RAKFRQIVSAVQYCHQKFIVHRDLKAENLLLDA
				DMNIKIADFGFSNEFTFGNKLDTFCGSPPYAAPEL
			1	FQGKKYDGPEVDVWSLGVILYTLVSGSLPFDGQ
L	L			NLKELRERVLRGKYRIPFYMSTDCENLLKKFLIL

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SEQ ID NO:	Method	Predicted beginning nucleotide location corresponding to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion
				NPSKRGTLEQIMKDRWMNVGHE\DDELKPYGEP LP\DYKDPRRTELMVSMGYTREEIQDSLVGQRYN EVMATYLLLGYKSSELEGDTITLKPRPSADLTNS SAPSPSHKVQRSVSANPKQRRFSDQAGPAIPTSNS YSKKTQSNNAENKRPEEDRESGRKASSTAKVPA SPLPGLERKKTTPTPSTNSVLSTSTNRSRNSPLL\E RASL\GQGFHPEWAKTALTMPGSRASTASASAA VSAARPRQHQKSMSASVHPNKASGLPPTESNCE VPRPRQVCWGSCTAPQRVPVASPSAHNISSSGGA PDRTNFPRGVSSRSTFHAGQLRQVR\DQQNLPYG VTPASPSGHSQGRRGASGSIFSKFTSKFVRRNLNE PESKDR\VETLRPHVV\NSGGNDKEKEEFREAKPR SLRFTWSMKTTSSMEPNEMMREIRKVLDANSCQ SELHEKYMLLCMHGTPGHEDFVQWEMEVCKLP RLSLNGVRFKRISGTSMAFKNIASKIANELKL
2980	A	120	3433	NCLLLQAKGFHGEIEDLQQWLTDTERHLLASKP LGGLPETAKEQLNVHMEVCAAFEAKEETYKSLM QKGQQMLARCPKSAETNIDQDINNLKEKWESVE TKLNER\KT\KLEEALNLA\MEFHNSL\QDFINWLT QAEQTLNVASRPSLILDTVLFQIDEHKVFANEVN SHREQIIELDKTGTHLKYFSQKQDVVLIKNLLISV QSRWEKVVQRLVERGRSLDDARKRAKQFHEAW SKLMEWLEESEKSLDSELEIANDPDKIKTQLAQH KEFQKSLGAKHSVYDTTNRTGRSLKEKTSLADD NLKLDDMLSELRDKWDTICGKSVERQNKLEEA\ LLFSGQFTDALQALIDWLYRVEPQLAEDQPVHG DIDLVMNLIDNHKAFQKELGKRTSSVQALKRSA RELIEGSRDDSSWVKVQMQELSTRWETVCALSIS KQTRLEAALRQAEEFHSVVHALLEWLAEAEQTL RFHGVLPDDEDALRTLIDQHKEFMKKLEEKRAE LNKATTMGDTVLAICHPDSITTIKHWITIIRARFEE VLAWAKQHQQRLASALAGLIAKQELLEALLAW LQWAETTLTDKDKEVIPQEIEEVKALIAEHQTFM EEMTRKQPDVDKVTKTYKRRAADPSSLQSHIPV LDKGRAGRKRFPASSLYPSGSQTQIETKNPRVNL LVSKWQQVWLLALERRRKLNDALDRLEELREF ANFDFDIWRKKYMRWMNHKKSRVMDFFRRIDK DQDGKITRQEFIDGILSSKFPTSRLEMSAVADIFD RDGDGYIDYYEFVAALHPNKDAYKPITDADKIE DEVTRQVAKCKCAKRFQVEQIGDNKYRFFLGNQ FGDSQQLRLVRILRSTVMVRVGGGWMALDEFL VKNDPCRAKGRTNMELREKFILADGASQGMAA FRPRGRRSRPSSRGASPNRSTSVSSQAAQAASPQ VPATTTPKILHPLTRNYGKPWLTNSKMSTPCKAA ECSDFPVPSAEGTPIQGSKLRLPGYLSGKGFHSGE DSGLITTAAARVRTQFADSKKTPSRPGSRAGSKA GSRASSRRGSDASDFDISEIQSVCSDVETVPQTHR PTPRAGSRPSTAKPSKIPTPQRKSPASKLDKSSKR
2981	A	120	3433	NCLLLQAKGFHGEIEDLQQWLTDTERHLLASKP LGGLPETAKEQLNVHMEVCAAFEAKEETYKSLM QKGQQMLARCPKSAETNIDQDINNLKEKWESVE TKLNER\KT\KLEEALNLA\MEFHNSL\QDFINWLT QAEQTLNVASRPSLILDTVLFQIDEHKVFANEVN SHREQIIELDKTGTHLKYFSQKQDVVLIKNLLISV QSRWEKVVQRLVERGRSLDDARKRAKQFHEAW

SEQ ID NO:	Method	Predicted beginning nucleotide location corresponding to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion
				SKLMEWLESEKSLDSELEIANDPDKIKTQLAQH KEFQKSLGAKHSVYDTTNRTGRSLKEKTSLADD NLKLDDMLSELRDKWDTICGKSVERQNKLEEA\ LLFSGQFTDALQALIDWLYRVEPQLAEDQPVHG DIDLVMNLIDNHKAFQKELGKRTSSVQALKRSA RELIEGSRDDSSWVKVQMQELSTRWETVCALSIS KQTRLEAALRQAEEFHSVVHALLEWLAEAEQTL RFHGVLPDDEDALRTLIDQHKEFMKKLEEKRAE LNKATTMGDTVLAICHPDSITTIKHWITIIRARFEE VLAWAKQHQQRLASALAGLIAKQELLEALLAW LQWAETTLTDKDKEVIPQEIEEVKALIAEHQTFM EEMTRKQPDVDKVTKTYKRRAADPSSLQSHIPV LDKGRAGRKRFPASSLYPSGSQTQIETKNPRVNL LVSKWQQVWLLALERRRKLNDALDRLEELREF ANFDFDIWRKKYMRWMNHKKSRVMDFFRRIDK DQDGKITRQEFIDGILSSKFPTSRLEMSAVADIFD RDGDGYIDYYEFVAALHPNKDAYKPITDADKIE DEVTRQVAKCKCAKRFQVEQIGDNKYRFFLGNQ FGDSQQLRLVRILRSTVMVRVGGGWMALDEFL VKNDPCRAKGRTNMELREKFILADGASQGMAA FRPRGRRSRPSSRGASPNRSTSVSSQAAQAASPQ VPATTTPKILHPLTRNYGKPWLTNSKMSTPCKAA ECSDFPVPSAEGTPIQGSKLRLPGYLSGKGFHSGE DSGLITTAAARVRTQFADSKKTPSRPGSRAGSKA GSRASSRRGSDASDFDISEIQSVCSDVETVPQTHR
2982	A	1	2065	PTPRAGSRPSTAKPSKIPTPQRKSPASKLDKSSKR MAAGGAEGGSGPGAAMGDCAEIKSQFRTREGF YKLLPGDGAARRSGPASAQTPVPPQPPQPPPGPA SASGPGAAGPASSPPPAGPGPGPALPAVRLSLVR LGEPDSAGAGEPPATPAGLGSGGDRVCFNLGRE LYFYPGCCRRGSQRWHTPLTPFLPPLKSIDLNKPI DKRIYKGTQPTCHDFNQFTAATETISLLVGFSAG QVQYLDLIKKDTSKLFNEERLIDKTKVTYLKWLP ESESLFLASHASGHLYLYNVSHPCASAPPQYSLL KQ\AWGFSFYAAKSKAPRNPLAKWAVGEGPLNE FAFSPDGRHLACVSQDGCLRVFHFDSMLLRGLM KSYFGGLLCVCWSPDGRYVVTGGEDDLVTVWS FTEGRVVARGHGHKSWVNAVAFDPYTTRAEEA ATAAGADGERSGEEEEEPEAAGTGSAGGAPLSP LPKAGSITYRFGSAGQDTQFCLWDLTEDVLYPHP PLARTRTLPGTPGTTPPAASSSRGGEPGPGPLPRS LSRSNSLPHPAGGGKAGGPGVAAEPGTPFSIGRF ATLTLQERRDRGAEKEHKRYHSLGNISRGGSGG SGSGGEKPSGPVPRSRLDPAKVLGTALCPRIHEV PLLEPLVCKKIAQERLTVLLFLEDCIITACQEGLIC TWARPGKAFTDEETEAQTGEGSWPRSPSKSVVE GISSQPGNSPSGTVV
2983	A	3855	220	RRFRLSAHRAQPCCRCRGLEMPRGVFQQLSNLV LQELNANLSNLTSAFEKATAEKIKCQQEADATN RVILLANRLVGGLASENIRWAESVENFRSQGVTL CGDVLLISAFVSYVGYFTKKYRNELMEKFWIPYI HNLKVPIPITNGLDPLSLLTDDADVATWNNQGLP SDRMSTENATILGNTERWPLIVDAQLQGIKWIKN KYRSELKAIRLGQKSYLDVIEQATSEGDTLLIENI GETVDPALDPLLGRNTIKKGKYIKIGDKEVGVPP

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SEQ ID NO:	Method	Predicted beginning nucleotide location corresponding to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion
	•			QVPPDPTHQVLQPTLQARDAGSVH\LINFLVTRD GLEDQLLAAVVAKERPDLEQLKANLTKSQNEFK IVLKELEDSLLARLSAASGNFLGDTALVENLETT KHTASEIEEKVVEAKITEVKINEARENYRPAAER ASLLYFILNDLNKINPVYQFSLKAFNVVFEKAIQR TTPANEVKQRVINLTDEITYSVYMYTARGLFERD KLIFLAQVTFQVLSMKKELNPVELDFLLRFPFKA GVVSPVDFLQHQGWGGIKALSEMDEFKNLDSDI EGSAKRWKKLVESEAPEKEIFPKEWKNKTALQK LCMVRCLRPDRMTYAIKNFVEEKMGSKFVEGRS VEFSKSYEESSPSTSIFFILSPGVDPLKDVEALGKK LGFTIDNGKLHNVSLGQGQEVVAENALDVAAEK GHWVILQNIHLVARWLGTLDKKLERYSTGRHED YRVFIRAEPAPSPETHIIPQGILENAIKITNEPPTGM YANLYKALDLFTQDTLEMCTKEMEFKCMLFAL CYFHAVVAERRKFGAQGWNRSYPFNNGDLTISI NVLYNYLEANPKVPWDDLRYLFGEIMYGGHITD DWDRRLCRTYLAEYIRTEMLEGDVLLAPGFQIPP NLDYKGYHEYIDENLPPESPYLYGLHPNAEIGFL TVTSEKLFRTVLEMQPKETDSGAGTGVSREEKV KAVLDDILEKIPETFNMAEIMAKAAEKTPYVVV AFQECERMNILTNEMRRSLKELNLGLKGELTITT DVEDLSTALFYDTVPDTWVARAYPSMMGLAAW YANLLLRIRELEAWTTDFALPTTVWLAGFFNPQS FLTAIMQSMARKNEWPLDKMCLSVEVTKKNRE DMTAPPREGSYVYGLFMEGARWDTQTGVIAEA RLKELTPAMPVIFIKAIPVARMETKNIYECPVYKT RIRGPTYVWTFNLKTKEKAAKWILAAVALLLQV
2984	A	2	1464	FVLFPGIAMETPGASASSLLLPAASRPPRKREAGE AGAATSKQRVLDEEEYIEGLQTVIQRDFFPDVEK LQAQKEYLEAEENGDLERMRQIAIKFGSALGKM SREPPPPYVTPATFETPEVHAGTGVVGNKPRPRG RGLEDGEAGEEEEKEPLPSLDVFLSRYTSEDNAS FQEIMEVAKERSRARHAWLYQAEEEFEKRQKDN LELPSAEHQAIESSQASVETWKYKAKNSLMYYP EGVPDEEQLFKKPRQVVHKNTRFLRDPFSQALSR CQLQQAAALNAQHKQGKVGPDGKELIPQESPRV GGFGFVATPSPAPGVNESPMMTWGEVENTPLRV EGSETPYVDRTPGPAFKILEPGRRERLGLKMANE AAAKNRAKKQEALRRVTENLASLTPKGLSPAMS PALQRLVSRTASKYTDRALRASYTPSPARSTHLK NPGPVGCRPPQSTPGA/PGSATRTPL\TQDPA\SIT DNLLQLPARRKASDFF
2985	A	1890	178	ASTQEAGLLSPPGVGAQRCWNFVACLPVRACAD MASNDYTQQATQSYGAQRCWNFVACLPVRACAD MASNDYTQQATQSYGAQRCWNFVACLPVRACAD YGQQSYSQYSQATQSYGAQSYSGYGQSYSGYGQSYSGYGQSYSGYGQSYGQYGSTGGYGSSGSYGQPQSSYPGY GQQPAPSSTSGSYGSSSQSSSYGQPQSGSYSQQPS YGGQQQSYGQQQSYNPPRGYGQQNQYNSSSGG GGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGG

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SEQ ID NO:	Method	Predicted beginning nucleotide location corresponding to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methlonine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion
				WFDGKEFSGNPIKVSFATRRADFNRGGGNGRGG RGRGGPMGRGGYGGGGSGGGGRGGFPSGGGGG GGQQRAGDWKCPNPTCENMNFSWRNECNQCK APKPDGPGGGPGGSHMGGNYGDDRRGGRGGYD RGGYRGRGGDRGGFRGGRGGDRGGFGPGKM DSRGEHRQDRRERPY
2986		1890	178	ASTQEAGLLSPPGVGAQRCWNFVACLPVRACAD MASNDYTQATQSYGAYPTQPGQGYSQQSSQP YGQQSYSGYSQSTDTSGYGQSSYSSYGQSQNSY GTQSTPQGYGSTGGYGSSQSSQSSYGQQSSYPGY GQQPAPSSTSGSYGSSSQSSSYGQPQSGSYSQQPS YGGQQQSYGQQQSYNPPRGYGQQNQYNSSSGG GGGGGGGGSYGQDQSSMSGSGGGGGGGGGGS GGGGGYGNQDQTGAAGSRGYRQ\QDRGGRCRG GSGGGGS\GGAAGYNRSSGGYEPRGRGGGRGGR GGMGGSDRGGFNKFGGPRDQGSRHDSEQDNSD NNTIFVQGLGENVTIESVADYFKQIGIIKTNKKTG QPMINLYTDRETGKLKGEATVSFDDPPSAKAAID WFDGKEFSGNPIKVSFATRRADFNRGGGNGRGG RGRGGPMGRGGYGGGGGGGGGGGGGGGGGGGGGGGGGGGG
2987	A	1376	898	GGAKAGGAPHPFTLPFRHVGGLSAAPEEVEGML WAGARQHGRNWRKRETSPGTQGPLPPVPR/VPP GPDG\PHAIAPTLSWAIPRQQCSPQPGRLNALPPD RCSGPHFGDRAPESCFPGACSVSGACAFKGTRPA CPPQEPSLRSSRNRLREGQTFGRMEI
2988	A	1		MGNDSVSYEYGDYSDLSDRPVDCLDGACLAIDP LRVAPLPLYAAIFLVGVPGNAMVAWVAGKVAR RRVGATWLLHLAVADLLCCLSLPILAVPIARGGH WPYGAVGCRALPSIILLTMYASVLLLAALSADLC FLALGPAW\CLRFS\GACGVQVACGAAWTLALL LTVPSAIYRRLHQEHFPARLQCVVDYGGSSSTEN AVTAIRFLFGFLGPLVAVASCHSALLCWAARRC RPLGTAIVVGFFVCWAPYHLLGLVLTVAAPNSA LLARALRAEPLIVGLALAHSCLNPMLFLYFGRAQ LRRSLPAACHWALRESQGQDESVDSKKSTSHDL VSEMEV
2989	A	27	4074	KSQLFCFWVGKAGDILSGDQDKEQKDPYFVETP YGYQLDLDFLKYVDDIQKGNTIKRLNIQKRRKPS VPCPEPRTTSGQQGIWTSTESLSSSNSDDNKQCP NFLIARSQVTSTPISKPPPPLETSLPFLTIPENRQLP PPSPQLPKHNLHVTKTLMETRRLEQERATMQM TPGEFRRPRLASFGGMGTTSSLPSFVGSGNHNPA KHQLQNGYQGNGDYGSYAPAAPTTSSMGSSIRH SPLSSGISTPVTNVSPMHLQHIREQMAIALKRLKE LEEQVRTIPVLQVKISVLQEEKRQLVSQLKNQRA ASQINVCGVRKRSYSAGNASQLEQLSRARRSGG ELYIDYEEEEMETVEQSTQRIKEFRQL\TADMQA LEQKIQDSSCEASSELRENGECRSVAVGAEENMN DIVVYHRGSRSCKDAAVGTLVEMRNCGVSVTEA MLGVMTEADKEIELQQQTIESLKEKIYRLEVQLR ETTHDREMTKLKQELQAAGSRKKVDKATMAQP

SEQ ID NO:	Method	Predicted beginning nucleotide location corresponding to first amino acid residue of peptide	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion
		sequence		LVFSKVVEAVVQTRDQMVGSHMDLVDTCVGTS VETNSVGISCQPECKNKVVGPELPMNWWIVKER VEMHDRCAGRSVEMCDKSVSVEVSVCETGSNTE ESVNDLTLLKTNLNLKEVRSIGCGDCSVDVTVCS PKECASRGVNTEAVSQVEAAVMAVPRTADQDT STDLEQVHQFTNTETATLIESCTNTCLSTLDKQTS TQTVETRTVAVGEGRVKDINSSTKTRSIGVGTLL SGHSGFDRPSAVKTKESGVGQININDNYLVGLK MRTIACGPPQLTVGLTASRRSVGVGDDPVGESLE NPQPQAPLGMMTGLDHYIERIQKLLAEQQTLLA ENYSELAEAFGEPHSQMGSLNSQLISTLSSINSVM KSASTEELRNPDFQKTSLGKITGSYLGYTCKCGG LQSGSPLSSQTSQPEQEVGTSEGKPISSLDAFPTQ EGTLSPVNLTDDQIAAGLYACTNNESTLKSIMKK KDGNKDSNGAKKNLQFVGINGGYETTSSDDSSS DESSSSESDDECDVIEYPLEEEEEEEDEDTRGMAE GHHAVNIEGLKSARVEDEMQVQECEPEKVEIRE RYELSEKMLSACNLLKNTINDPKALTSKDMRFC LNTLQHEWFRVSSQKSAIPAMVGDYIAAFEAISP DVLRYVINLADGNGNTALHYSVSHSNFEIVKLLL DADVCNVDHQNKAGYTPIMLAALAAVEAEKDM RIVEELFGCGDVNAKASQAGQTALMLAVSHGRI DMVKGLLACGADVNIQDDEGSTALMCASEHGH VEIVKLLLAQPGCNGHLEDNDGSTALSIALEAGH KDÏAVLLYAHVNFAKAQSPGTPRLGRKTSPGPTH
2990	A		1687	RGSFD ERLRPGQRAIRGPVPAAGACASLPPRAGPAQGRH AALGGAEPGSHLHCGVRLQRREEPGGQQRLLPQ RGGSAQTGHQHPGPYECQCPGPQPGGTTPALLSL ILEETRGPPASANPDKDHSTQPGTMGRKKIQISRI LDQRNRQVTFTKRKFGLMKKAYELSVLCDCEIA LIIFNSATRLFQYASTDMDRVLLKYTEYSEPHESR TNTDILETLKRRGIGLDGPELEPDEGPEEPGEKFR RLAGEGGDPALPRPRLYPAAPAMPSPDVVYGAL PPPG\CDPSGLGEALPAQSRPSPFRPAAPKAGPPG LGHPLFSPSHLTSKTPPPLYLPTEGRRSDLPGGLA GPRGGLNTSRSLYSGLQNPCSTATPGPPLGSFPFL PGGPPVGAEAWARRVPQPAAPPRRPPQSSIKSER LFLRPPGAPATFLRPSPIPCSSPGPWQSLCGLGPP\ CAGCPWPTAGPGRRSPGGTSPERSPGTARARGDP \TSLQAFSEKTHTVTAPLRGGGLEVGGWTQSSAG GLLSFFLFVCISTNKNARGVRGPEKK
2991	A	3	1159	IPQPLHCASPKEEMSLRCGDAARTLGPRVFGRYF CSPVRPLSSLPDKKKELLQNGPDLQDFVSGDLAD RSTWDEYKGNLKRQKGERLRLPPWLKTEIPMGK NYNKLKNTLRNLNLHTVCEEARCPNIGECWGGG EYATATATIMLMGDTCTRGCRFCSVKTARNPPP LDASEPYNTAKAIAEWGLDYVVLTSVDRDDMP DGGAEHIAKTVSYLKERNPKILVECLTPDFRGDL KAIEKVALSGLDVYAHNVETVPELQSKVRDPRA NFDQSLRVLKHAKKVQPDVISKTSIMLGLGENDE QVYATMKALREADVDCLTLGQYMQPTRRHLKV EEYITPEKFKYWEKVGNELGFHYTASGP\LVRSS YKAGEFFLKNLVAKRKTKDL
2992	Α	3	1636	PVPGVPTSPPSCCPQDMQGPWVLLLLGLRLQLSL

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				GVIPAEEENPAFWNRQAAEALDAAKKLQPIQKV AKNLILFLGDGLGVPTVTATRILKGQKNGKLGPE TPLAMDRFPYLALSKTYNVDRQVPDSAATATAY LCGVKANFQTIGLSAAARFNQCNTTRGNEVISV MNRAKQAGKSVGVVTTTRVQHASPAGTYAHTV NRNWYSDADMPASARQEGCQDIATQLISNMDID VILGGGRKYMFPMGTPDPEYPADASQNGIRLDG
				KNLVQEWLAKHQGAWYVWNRTELMQASLDQS VTHLMGLFEPGDTKYEIHRDPTLDPSLMEMTEA ALRLLSRNPRGFYLFVEGGRIDHGHHEGVAYQA LTEAVMFDDAIERAGQLTSEEDTLTLVTADHSH VFSFGGYTLRGSSIFGLAPSKAQDSKAYTSILYGN GPGYVFNSGVRPDVNESESGSPDYHQQAG\VPLS SETHGGEDVAVFARGPQAHLVHGVQEQSFVAH VMAFAACLEPYTACDLAPPACTTDAAHPVAASL PLLAGTLLLLGASAAP
2993	A	3	685	DAWARLKMNRLFGKAKPKAPPPSLTDCIGTVD SRAESIDKKISRLDAELVKYKDQIKKMREGPAKN MVKQKALRVLKQKRMYEQQRDNLA\NSHSTW\ TS\HYTIQSLKDTKTTVDAMKLGVKEMKKAYKQ VKIDQIEDLQDQLEDMMEDANEIQEALSRSYGTP ELDEDDLEAELDALGDELLADEDSSYLDEAASA PAIPEGVPTDTKNKDGVLVDEFGLPQIPAS
2994	A	1710	161	RRCELTPFIKTLILPKSWGAFPEDVVMQHVSSSQ SSQRHVQWPGACPGAGEEQPACSQPSLPLTLPSP SHQLQQLMVRGGPAGGQNMNVDLQGVGPGLQ GSPQVTLAPLPLPSPTSPGFQFSAQPRRFEHGSPS YIQVTSPLSQQVQTQSPTQPSPGPGQALQNVRAG APGPGLGLCSSSPTGDFVDASVLVRQISLSPSSGG HFVFQDGSGLTQIAQGAQVQLQHPGTPITVRERR PSQPHTQSGGTIHHLGPQSPAAAGGAGLQPLASP SHITTANLPPQISSIIQGQLVQQQQVLQGPPLPRPL GFERTPGVLLPGAGGAAGFGMTSPPPPTSPSRTA VPPGLSSLPLTSVGNTGMKKVPKKLEEIPPASPE MAQMRKQCLDYHHQEMQALKEVFKEYLIELFF LQHFQGNMMDFLAFKERLYGPLQAYLRQNDLDI EEEEEE\HFEVINDEVKVVARKHGQPGTPVAIAT\ QLPPRTSAAFPAQQQPLQVLSDGSTVQLPRLSSL GFEDSMC
2995	A	3	924	SAPSGIDASTHAFARCKHPINVRRDPSIPIYGLRQS ILLNTRLQDCYVDSPALTNIWMARTCAKQNINAP APATTSSWEVVRNPLIASSFSLVKLVLRRQLKNK CCPPPCKFGEGKLSKRLKHKDDSVMKATQQARK RNFISSKSKQPAGHRRPAGGIRESKESSKEKKLTV RQDLEDRYAEHVAAT\QALPQDSGTAAWKG\RV LLPETQKRQQLSEDTLTIHGLPTEGYQALYHAVV EPMLWNPSGTPKRYSLELGKAIKQKLWEALCSQ GAISEGAQRDRFPGRKQPGVHEEPVLKKWPKLK SKK
2996	Α .	3	1713	GKFGIKPSQRRISGKSTFHSEMEGEDTRDDSLYSI LEELWQDAEQIKRCQEKHNKLLSRTTFLNKKILN TEWDYEYKDFGKFVHPSPNLILSQKRPHKRDSFG KSFKHNLDLHIHNKSNAAKNLDKTIGHGQVFTQ NSSYSHHENTHTGVKFCERNQCGKVLSLKHSLS QNVKFPIGEKANTCTEFGKIFTQRSHFFAPQKIHT

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				VEKPHELSKCVNVFTQKPLLSIYLRVHRDEKLYI\ CTKM/CGKGLHPRNSELIMHEKTHTREKPYKCNE \CGKSFFQVSSLLRHQTTHTGEKLFECSECGKGFS LNSALNIHQKIHTGERHHKCSECGKAFTQKSTLR MHQRIHTGERSYICTQCGQAFIQKAHLIAHQRIH TGEKPYECSDCGKSFPSKSQLQMHKRIHTGEKPY ICTECGKAFTNRSNLNTHQKSHTGEKSYICAECG KAFTDRSNFNKHQTIHTGEKPYVCADCGRAFIQK SELITHQRIHTTEKPYKCPDCEKSFSKKPHLKVHQ RIHTGEKPYICAECGKAFTDRSNFNKHQTIHTGD KPYKCSDCGKGFTQKSVLSMHRNIHT
2997	A	3	1763	AASTRTMGSRHFEGIYDHVGHFGRFQRVLYFICA FQNISCGIHYLASVFMGVTPHHVCRPPGNVSQVV FHNHSNWSLEDTGALLSSGQKDYVTVQLQNGEI WELSRCSRNKRENTSSLGYEYTGSKKEFPCVDG YIYDQNTWKSTAVTQWNLVCDRKWLAMLIQPL FMFGGPTGIG/VTFGYF\SDRLGRRVVLWATSSS MFLFGIAAAFAVDYYTFMAARFFLAMVASGYLV VGFVYVMEFIGMKSRTWASVHLHSFFAVGTLLV ALTGYLVRTWWLYQMILSTVTVPFILCCWVLPE TPFWLLSEGRYEEAQK\IVDIMAKWNRASSCKLS ELLSLDLQGPVSNSPTEVQKHNLSYLFYNWSITK RTLTVWLIWFTGSLGFYSFSLNSVNLGGNEYLNL FLLGVVEIPAYTFVCIAMDKVGRRTVLAYSLFC\S ALACGVVMVIPQKHYILGVVTAM\VGKILPIGAA FG\LIYLYTAELYPTIVRSLAVGSGSMVCRLASIL APFSVDLSSIWIFIPQLFVGTMALLSGVLTLKLPE TLGKRLATTWEEAAKLESENESKSSKLLLTTNNS GLEKTEAITPRDSGLGE
2998	A	3	1441	QRPASQLLAPFAAEALPGAPRAAMAQHFSLAAC DVVGFDLDHTLCRYNLPESAPLIYNSFAQFLVKE KGYDKELLNVTPEDWDFCCKGLALDLEDGNFL KLANNGTVLRASHGTKMMTPEVLAEAYGKKEW KHFLSDTGMACRSGKYYFYDNYFDLPGALLCAR VVDYLTKLNNGQKTFDFWKDIVAAIQHNYKMS AFKENCGIYFPEIKRDPGRYLHSRPESVKKWLRQ LKNAGKILLLITSSHSDYCRLLCAYILGNDFTDLF DIVITNALKPGFFSHLPSQRPFRTLENDEEQEALP SLDKPGWYSQGNAVHLYELLKKMTGKPEPKVV YFGDSMHSDIFPARHYSNWETVLILEELRGDEGT RSQRPEESEPLEKKGKYEGPKAKPLNTSSKKWGS FF\IDSVLGLENTEDSLVYTWSCKRISTYSTIAIPSI EAIAELPLDYKFTRFSSSNSKTAGYYPNPPLVLSS DETLISK
2999	A	320	2417	LRRKMTPQSLLQTTLFLLSLLFLVQGAHGRGHR EDFRFCSQRNQTHRSSLHYKPTPDLRISIENSEEA LTVHAPFPAAHPASRSFPDPRGLYHFCLYWNRH AGRLHLLYGKRDFLLSDKASSLLCFQHQEESLAQ GPPLLATSVTSWWSPQNISLPSAASFTFSFHSPPH TGAHNASVDMCELKRDLQLLSQFLKHPQKASRR PSAAPASQQLQSLESKLTSVRFMGDMGSFEEDRI NATVWKLQPTAGLQDLHIHSRQEEEQSEIMEYS VLLPRTLFQRTKGRSGEAEKRLLLVDFSSQALFQ DKNSSQVLGEKVLGIVVQNTKVANLTEPVVLTF QHQLQPKNVTLQCVFWVEDPTLSSPGHWSSAGC

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SEQ ID NO:	Method	Predicted beginning nucleotide location corresponding to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion
				ETVRRETQTSCFCNHLTYFAVLMVSSVEVDAVH KHYLSLLSYVGCVVSALACLVTIAAYLCSRVPLP CRRKPRDYTIKVHMNLLLAVFLLDTSFLLSEPVA LTGSEAGCRASAIFLHFSLLTCLSWMGLEGYNLY RLVVEVFGTYVPGYLLKLSAMGWGFPIFLVTLV ALVDVDNYGPIILAVHRTPEGVIYPSMCWIRDSL VSYITNLGLFSLVFLFNMAMLATMVVQILRLRPH TQKWSHVLTLLCLSLVLG\LPWALIFFSFASGTFQ LVVLYLFSIITSFQGFLIFIWYWSMRLQARGGPSP LKSNSDSARLPISSGSTSSSRI
3000	A	66	1003	SRGQLDAGQSSEQHGGNRQPEQSRSRSSSSSSP RRSRSAAEPAMALSMPLNGLKEEDKEPLIELFVK AGSDGESIGNCPFSQRLFMILWLKGVVFSVTTVD LKRKPADLQNLAPGTHPPFITFNSEVKTDVNKIEE FLEEVLCPPKYLKLSPKHPESNTAGMDIFAKFSA YIKNSRPEANEALERGLLKTLQKLDEYLNSPLPD EIDENSMEDIKFSTRKFLDGNEMTLADCNLLPKL HIVKVVAKKYRNFDIPKEMTGIWRYLTNAYSRD EFTNTCPSDKEVEI\AYSDVAKRLHQVKSRLLKE VSFMSSP
3001	A	779	2006	LALTFRSALSTLPGSPMTSSGSPDLQLAWGPSLLP HPPSVWSPALPSCFAGPCPLLPLSDTQGWWGPN WLAPPSAALCRPDAAVWPDLPSSNILLVTPPPAK *SAVAV*PCPRGAHSLERAARQYTISGSSTSQSGK CSKRDTKCCAVTTSWGCFWQKHWKGDEDSGW AFQEGSHLGEGHL
3002	A	909	2799	VEEAWTVWLHWGVRECLLEETNQKEEAASSN WTKARGPFWQEDWVWDMRLKMTTRNFPEREV PCDVEVERFTREVPCLSSLGDGWDCENQEGHLR QSALTLEKPGTQEAICEYPGFGEHLIASSDLPPSQ RVLATNGFHAPDSNVSGLDCDPALPSYPKSYAD KRTGDSDACGKGFNHSMEVIHGRNPVREKPYKY PESVKSFNHFTSLGHQKIMKRGKKSYEGKNFENI FTLSSSLNENQRNLPGEKQYRCTECGKCFKRNSS LVLHHRTHTGEKPYTCNECGKSFSKNYNLIVHQ RIHTGEKPYECSKCGKAFSDGSALTQHQRIHTGE KPYECLECGKTFNRNSSLILHQRTHTGEKPYRCN ECGKPFTDISHLTVHLRIHTGEKPYECSKCGKAF RDGSYLTQHERTHTGEKPFECAECGKSFNRNSHL IVHQKIHSGEKPYECKECGKTFIESAYLIRHQRIH TGEKPYGCNQCQKLFRNIAGLIRHQRTHTGEKPY ECNQCGKAFRDSSCLTKHQRIHTKETPYQCPECG KSFKQNSHLAVHQRLHSREGPSRCPQCGKMFQK SSSLVRHQRAHLGEQPMET*WLGAT*VFQFTLTP VFRRVLDLTPLWSVEKNPLSYPVN
3003	A	2	1489	SLTEHLSFFQPTAHSLTSLLGTMTTCSRQFTSSSS MKGSCGIGGGIGGGSSRISSVLAGGSCRAPSTYG GGLSVSSRFSSGGACGLGGGYGGGFSSSSSFGSG FGGGYGGGLGAGFGGGLGAGFGGGFAGGDGLL VGSEKVTMQNLNDRLASYLDKVRALEEANADL EVKIRDWYQRQRPSEIKDYSPYFKTIEDLRNKIIA ATIENAQPILQIDNARLAADDFRTKYEHELALRQ TVEADVNGLRRVLDELTLARTDLEMQIEGLKEE LAYLRKNH*EEMLALRGQTGGEVNVETDAAPG VDLSCILNEMRNQYEQMAEKNRRDAETWFLSKT

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				EELNKEVASNSELVQSSRSEVTELRRVLQGLEIEL QSQLSMKASLENSLEETKGRYCMQLSQIQGLIGS VEEQLAQLRCEMEQQSQEYQILLDVKTRLEQEIA TYRRLLEGEDAHLSSQQASGQSYSSREVFTSSSSS SSRQTRPILKEQSSSSFSQGQSS
3004		2	940	GCAPDTRFFVPEPGGRGAAPWVALVARGGCTFK DKVLVAARRNASAVVLYNEERYGNITLPMSHAG TGNIVVIMISYPKGREILELVQKGIPVTMTIGVGT RHVQEFISGQSVVFVAIAFITMMIISLAWLIFYYIQ RFLYTGSQIGSQSHRKETKKVIGQLLLHTVKHGE KGIDVDAENCAVCIENFKVKDIIRILPCKHIFHRIC IDPWLLDHRTCPMCKLDVIKALGYWGEPGDVQE MPAPESPPGRDPAANLSLALPDDDGSDESSPPSA SPAESEPQCDPSFKGDAGENTALLEAGRSDSRHG GPIS
3005		184	2552	TMTIHQFLLLFLFWVCLPHFCSPEIMFRRTPVPQQ RILSSRVPRSDGKILHRQKRGWMWNQFFLLEEY TGSDYQYVGKLHSDQDKGDGSLKYILSGDGAGT LFIIDEKTGDIHATRRIDREEKAFYTLRAQAINRR TLRPVEPESEFVIKIHDINDNEPTFPEEIYTASVPE MSVVGTSVVQVTATDADDPSYGNSARVIYSILQ GQPYFSVEPETGIIRTALPNMNRENREQYQVVIQ AKDMGGQMGGLSGTTTVNITLTDVNDNPPRFPQ NTIHLRVLESSPVGTAIGSVKATDADTGKNAEVE YRIIDGDGTDMFDIVTEKDTQEGIITVKKPLDYES RRLYTLKVEAENTHVDPRFYYLGPFKDTTIVKISI EDVDEPPVFSRSSYLFEVHEDIEVGTIIGTVMARD PDSISSPIRFSLDRHTDLDRIFNIHSGNGSLYTSKP LDRELSQWHNLTVIAAEINNPKETTRVAVFVRIL DANDNAPQFAVFYDTFVCENARPGQLIQTISAVD KDDPLGGQKFFFSLAAVNPNFTVQDNEDNTARIL TRKNGFNRHEISTYLLPVVISDNDYPIQSSTGTLTI RVCACDSQGNMQSCSAEALLLPAGLSTGALIAIL LCIIILLVIVVLFAALKRQRKKEPLILSKEDIRDNIV SYNDEGGGEEDTQAFDIGTLRNPAAIEEKKLRRD IIPETLFIPRRTPTAPDNTDVRDFINERLKEHDLDP TAPPYDSLATYAYEGNDSIAESLSSLESGTTEGD QNYDYLREWGPRFNKLPQKYGGGESDKDS
3006	A	2	541	GRVDKTWWGKSVGIMLTELEKALNSIIDVYHKY SLIKGNFHAVYRDDLKKLLETECPQYIRKKGAD VWFKELDINTDGAVNFQEFLILVIKMGVAALNSII DVYHKYSLIKGNFHAVYRDDLQKLLETECPQYI RKKGADVWFKELDINTDGAVNFQEFLILVIKMG VGSPQKKVASYF
3007	A	1	1253	MYEGIRCLLKALLGFVSLAIGTLYCPRQYRPFPG SLGIEAINVPEPIPDSYYRDMATWPTHAPSVEEG GQGRFGNQADHFLGSLAFAKLLNRSLAVPSWIE YQHHKPPFTNLHVSYQKYFKLEPLQAYHRVISLE DFMEKLAPTHWPPEKRVAYCFEVAAQRSPDKKT CPMKEGNPFGPFWDQFHVSFNKSELFTGISFSAS YREQWSQRFSPKEHPVLALPGAPAQFPVLEEHRP LQKYMVWSDEMVKTGEAQIHAHLVRPYVGIHL RIGSDWKNACAMLKDGTAGSHFMASPQCVGYS RSTAAPLTMTMCLPDLKEIQRAVKLWVRSLDAQ SVYVATDSESYVPELQQLFKGKVKVVSLKPEVA

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				QVDLYILGQADHFIGNCVSSFTAFVKRERDLQGR PSSFFGMDRPPKLRDEF
3008	A	3136	1898	TARGGSEPGPTMAANYSSTSTRREHVKVKTSS QPGFLERLSETSGGMFVGLMAFLLSFYLIFTNEG RALKTATSLAEGLSLVVSPDSIHSVAPENEGRLV HIIGALRTSKLLSDPNYGVHLPAVKLRRHVEMY QWVETEESREYTEDGQVKKETRYSYNTEWRSEII NSKNFDREIGHKNPRAMAGESFMATAPFVQIGRF FLSSGLIDKVDNFKSLSLSKLEDPHVDIIRRGDFF YHSENPKYPEVGDLRVSFSYAGLSGDDPDLGPA HVVTVIARQRGDQLVPFSTKSGDTLLLHHGDFS AEEVFHRELRSNSMKTWGLRAAGWMAMFMGL NLMTRILYTLVDWFPVFRDLVNIGLKAFAFCVAT SLTLLTVAAGWLFYRPLWALLIAGLALVPILVAR TRVPAKKLE
3009	A	93	659	DAAVAMTAQGGLVANRGRRFKWAIELSGPGGG SRGRSDRGSGQGDSLYPVGYLDKQVPDTSVQET DRILVEKRCWDIALGPLKQIPMNLFIMYMAGNTI SIFPTMMVCMMAWRPIQALMAISATFKMLESSS QKFLQGLVYLIGNLMGLALAVYKCQSMGLLPTH ASDWLAFIEPPERMEFSGGGLLL
3010	A .	2		LIDSAKARYWTQRGTWVYDNALLLLLKCLWSN VVPECTMASSNTVLMRLVASAYSIAQKAGMIVR RVIAEGDLGIVEKTCATDLQTKADRLAQMSICSS LARKFPKLTIIGEEDLPSEEVDQELIEDSQWEEILK QPCPSQYSAIKEEDLVVWVDPLDGTKEYTEGLL DNVTVLIGIAYEGKAIAGVINQPYYNYEAGPDAV LGRTIWGVLGLGAFGFQLKEVPAGKHIITTTRSH SNKLVTDCVAAMNPDAVLRVGGAGNKIIQLIEG KASAYVFASPGCKKWDTCAPEVILHAVGGKLTD IHGNVLQYHKDVKHMNSAGVLATLRNYDYYAS RVPESIKNALVP
3011	A	291	1452	SPQKTMRSHTITMTTTSVSSWPYSSHRMRFITNH SDQPPQNFSATPNVTTCPMDEKLLSTVLTTSYSVI FIVGLVGNIIALYVFLGIHRKRNSIQIYLLNVAIAD LLLIFCLPFRIMYHINQNKWTLGVILCKVVGTLFY MNMYISIILLGFISLDRYIKINRSIQQRKAITTKQSI YVCCIVWMLALGGFLTMIILTLKKGGHNSTMCF HYRDKHNAKGEAIFNFILVVMFWLIFLLIILSYIKI GKNLLRISKRRSKFPNSGKYATTARNSFIVLIIFTI CFVPYHAFRFIYISSQLNVSSCYWKEIVHKTNEIM LVLSSFNSCLDPVMYFLMSSNIRKIMCQLLFRF QGEPSRSESTSEFKPGYSLHDTSVAVKIQSSSKST
3012	A	246	379	TEPVGYTKAEEPIAMRSLGALLLLLSACLAVSAG PVPTPPDNIQVQENFNISRIYGKWYNLAIGSTCPW LKKIMDRMTVSTLVLGEGATEAEISMTSTRWRK GVCEETSGAYEKTDTDGKFLYHKSKWNITMESY VVHTNYDEYAIFLTKKFSRHHGPTITAKLYGRAP QLRETLLQDFRVVAQGVGIPEDSIFTMADRGECV PGEQEPEPILIPRVRRAVLPQEEEGSGGGQLVTEV TKKEDSCQLGYSAGPCMGMTSRYFYNGTSMAC ETFQYGGCMGNGNNFVTEKECLQTCRTVAACN LPIVRGPCRAFIQLWAFDAVKGKCVLFPYGGCQ GNGNKFYSEKECREYCGVPGDGDEELLRFSN RQMALLKANKDLISAGLKEFSVLLNQQVFNDPL

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				VSEEDMVTVVEDWMNFYINYYRQQVTGEPQER DKALQELRQELNTLANPFLAKYRDFLKSHELPSH PPPSS
3014	A	1	373	GTSWSTLRAVMSASVVSVVSRVLEEYLSSTPQRL KLLDAYLLYILLTGALQFGYCLFVLTFHFNSLLLF FFFCVGSFHSNVYFLLFTLSFLCFLFIAYFFLIRFFS LFIWFFHVFFIELSLFYF
3015	A	2	1321	AAAEGTAPSPGRVSPPTPARGEPEVTVEIGETYLC RRPDSTWHSAEVIQSRVNDQEGREEFYVHYVGF NRRLDEWVDKNRLALTKTVKDAVQKNSEKYLS ELAEQPERKITRNQKRKHDEINHVQKTYAEMDP TTAALEKEHEAITKVKYVDKIHIGNYEIDAWYFS PFPEDYGKQPKLWLCEYCLKYMKYEKSYRFHLG QCQWRQPPGKEIYRKSNISVYEVDGKDHKIYCQ NLCLLAKLFLDHKTLYFDVEPFVFYILTEVDRQG AHIVGYFSKEKESPDGNNVACILTLPPYQRRGYG KFLIAFSYELSKLESTVGSPEKPLSDLGKLSYRSY WSWVLLEILRDFRGTLSIKDLSQMTSITQNDIIST LQSLNMVKYWKGQHVICVTPKLVEEHLKSAQY KKPPITGGWGAAVCRGRWGSVSIWTGRSQGLLI AVT
3016	A	2		AAAEGTAPSPGRVSPPTPARGEPEVTVEIGETYLC RRPDSTWHSAEVIQSRVNDQEGREEFYVHYVGF NRRLDEWVDKNRLALTKTVKDAVQKNSEKYLS ELAEQPERKITRNQKRKHDEINHVQKTYAEMDP TTAALEKEHEAITKVKYVDKIHIGNYEIDAWYFS PFPEDYGKQPKLWLCEYCLKYMKYEKSYRFHLG QCQWRQPPGKEIYRKSNISVYEVDGKDHKIYCQ NLCLLAKLFLDHKTLYFDVEPFVFYILTEVDRQG AHIVGYFSKEKESPDGNNVACILTLPPYQRRGYG KFLIAFSYELSKLESTVGSPEKPLSDLGKLSYRSY WSWVLLEILRDFRGTLSIKDLSQMTSITQNDIIST LQSLNMVKYWKGQHVICVTPKLVEEHLKSAQY KKPPITGGWGAAVCRGRWGSVSIWTGRSQGLLI AVT
3017	A	38	704	EAHPGGQLGSERNGVRMDEDVLTTLKILIGESG VGKSSLLLRFTDDTFDPELAATIGVDFKVKTISVD GNKAKLAIWDTAGQERFRTLTPSYYRGAQGVIL VYDVTRRDTFVKLDNWLNELETYCTRNDIVNM LVGNKIDKENREVDRNEGLKFARKHSMLFIEAS AKTCDGVQCAFEELVEKIIQTPGLWESENQNKG VKLSHREEGQGGGACGGYCSVL
3018	A	2640	2861	APVLILQMVKLSIVLTPQFLSHDQGQLTKELQQH VKSVTCPCEYLRKVSECRQMGPGALEQFPGLSC HTSHSG
3019	A	1307	711	PGITMAASLVGKKIVFVTGNAKKLEEVVQILGDK FPCTLVAQKIDLPEYQGEPDEISIQKCQEAVRQV QGPVLVEDTCLCFNALGGLPGPYIKWFLEKLKPE GLHQLLAGFEDKSAYALCTFALSTGDPSQPVRLF RGRTSGRIVAPRGCQDFGWDPCFQPDGYEQTYA EMPKAEKNAVSHRFRALLELQEYFGSLAA
3020	A	1202	180	VSCLPTSCKMITLNNQDQPVPFNSSHPDEYKIAA LVFYSCIFIIGLFVNITALWVFSCTTKKRTTVTIYM MNVALVDLIFIMTLPFRMFYYAKDEWPFGEYFC QILGALTVFYPSIALWLLAFISADRYMAIVQPKY

SEQ ID	Method	Predicted	Predicted end	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid,
NO:]	beginning	nucleotide	E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine,
1		nucleotide	location	I=Isoleucine, K=Lysine, L=Leucine, M=Methionine,
	1	location	corresponding	N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine,
[corresponding	to last amino	T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine,
:	1	to first amino	acid residue of	X=Unknown, *=Stop codon, /=possible nucleotide deletion,
	ļ	acid residue of	peptide	\=possible nucleotide insertion
]	1	peptide sequence	sequence	
	 	sequence		AKELKNTCKAVLACVGVWIMTLTTTTPLLLLYK
]	ļ		}	DPDKDSTPATCLKISDIIYLKAVNVLNLTRLTFFF
	į	i .		LIPLFIMIGCYLVIIHNLLHGRTSKLKPKVKEKSIRI
!	1		ĺ	IITLLVQVLVCFMPFHICFAFLMLGTGENSYNPW
	-			GAFTTFLMNLSTCLDVILYYIVSKQFQARVISVM
ļ		· ·		LYRNYLRSMRRKSFRSGSLRSLSNINSEML
2021	 	27	1897	EEFCTWIAVRVGEMETAPKPGKDVPPKKDKLQT
3021	Α	27	1097	-
1	ĺ			KRKKPRRYWEEETVPTTAGASPGPPRNKKNREL
ļ	Ì			RPQRPKNAYILKKSRISKKPQVPKKPREWKNPES
i	ļ			QRGLSGAQDPFPGPAPVPVEVVQKFCRIDKSRKL
	ļ		į	PHSKAKTRSRLEVAEAEEEETSIKAARSELLLAEE
l	1			PGFLEGEDGEDTAKICQADIVEAVDIASAAKHFD
1				LNLRQFGPYRLNYSRTGRHLAFGGRRGHVAALD
	1		1 : .	1
1				WYTKKLMCEINVMEAVRDIRFLHSEALLAVAQN
	1			RWLHIYDNQGIELHCIRRCDRVTRLEFLPFHFLLA
		ļ		TASETGFLTYLDVSVGKIVAALNARAGRLDVMS
	ł			QNPYNAVIHLGHSNGTVSLWSPAMKEPLAKILC
		·		HRGGVRAVAVDSTGTYMATSGLDHQLKIFDLRG
			1	TYQPLSTRTLPHGAGHLAFSQRGLLVAGMGDVV
ļ	ł	1		NIWAGQGKASPPSLEQPYLTHRLSGPVHGLQFCP
				FEDVLGVGHTGGITSMLVPGAGEPNFDGLESNPY
				RSRKQRQEWEVKALLEKVPAELICLDPRALAEV
i				DVISLEQGKKEQIERLGYDPQAKAPFQPKPKQKG
	Ĭ		ļ	RSSTASLVKRKRKVMDEEHRDKVRQSLQQQHH
	i			KEAKAKPTGARPSALDRFVR
3022	A	1	2249	MTAQDSNTSAHAQRDGPELPASSSWRSFWPLSC
3022	A	1	2249	
				LSSPPVSAVEVATEGRDREVAKVGQRFCDTTSGE
	ł			LRQARDRDCCVRMPAPVGRRSPPSPRSSMAAVA
	İ	j		LRDSAQGMTFEDVAIYFSQEEWELLDESQRFLYC
	ĺ	-	[DVMLENFAHVTSLGYCHGMENEAIASEQSVSIQ
	ļ.			VRTSKGNTPTQKTHLSEIKMCVPVLKDILPAAEH
1	į.			QTTSPVQKSYLGSTSMRGFCFSADLHQHQKHYN
	ľ			EEEPWKRKVDEATFVTGCRFHVLNYFTCGEAFP
			1	APTDLLQHEATPSGEEPHSSSSKHIQAFFNAKSYY
	1			KWGEYRKASSHKHTLVQHQSVCSEGGLYECSK
		1	1	CEKAFTCKNTLVQHQQIHTGQKMFECSECEESFS
				KKCHLILHKIIHTGERPYECSDREKAFIHKSEFIHH
	1			ORRHTGGVRHECGECRKTFSYKSNLIEHQRVHT
]			[GERPYECGECGKSFRQSSSLFRHQRVHSGERPYQ
				CCECGKSFRQIFNLIRHRRVHTGEMPYQCSDCGK
	1		1	SFSCKSELIQHQRIHSGERPYECRECGKSFRQFSN
	1			LIRHRSIHTGDRPYECSECEKSFSRKFILIQHQRVH
				TGERPYECSECGKSFTRKSDLIQHRRIHTGTRPYE
			1	CSECGKSFRQRSGLIQHRRLHTGERPYECSECGK
			1	SFSQSASLIQHQRVHTGERPYQCCECGKSFRQIFN
	1	ĺ	1	LIRHRRVHTGEMPYQCSDCGKSFSCKSELIQHRRI
1	[1	1	HSGERPYECSECGKSFSRKSNLIRHRRVHTEERP
	<u> </u>	L	-	
3023	A	3148	634	AAGALRCLAAFPRAEPASRGRQSSPARACAASR
			Į.	AERATAAAMAHRCLRLWGRGGCWPRGLQQLL
1	1			VPGGVGPGEQPCLRTLYRFVTTQARASRNSLLTD
	1		1	IIAAYQRFCSRPPKGFGKYFPNGKNGKKASEPKE
,				VMGEKKESKPAATTRSSGGGGGGGGKRGGKKD
	1	}	1	DSHWWSRFQKGDIPWDDKDFRMFFLWTALFWG
			1	
			1	GVMFYLLLKRSGREITWKDFVNNYLSKGVVDRL
1		<u> </u>	<u> </u>	EVVNKRFVRVTFTPGKTPVDGQYVWFNIGSVDT

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SEQ ID NO:	Method	Predicted beginning nucleotide location corresponding to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion
				FERNLETLQELGIEGENRVPVVYIAESDGSFLLS MLPTVLIIAFLLYTIRRGPAAIGRTGRGMGGLFSV GETTAKVLKDEIDVKFKDVAGCEEAKLEIMEFV NFLKNPKQYQDLGAKIPKGAILTGPPGTGKTLLA KATAGEANVPFITVSGSEFLEMFVGVGPARVRDL FALARKNAPCILFIDEIDAVGRKRGRGNFGGQSE QENTLNQLLVEMDGFNTTTNVVILAGTNRPDILD PALLRPGRFDRQIFIGPPDIKGRASIFKVHLRPLKL DSTLEKDKLARKLASLTPGFSGADVANVCNEAA LIAARHLSDSINQKHFEQAIERVIGGLEKKTQVLQ PEEKKTVAYHEAGHAVAGWYLEHADPLLKVSII PRGKGLGYAQYLPKEQYLYTKEQLLDRMCMTL GGRVSEEIFFGRITTGAQDDLRKVTQSAYAQIVQ FGMNEKVGQISFDLPRQGDMVLEKPYSEATARLI DDEVRILINDAYKRTVALLTEKKADVEKVALLL LEKEVLDKNDMVELLGPRPFAEKSTYEEFVEGT GSLDEDTSLPEGLKDWNKEREKEKEEPPGEKVA N
3024	A	274	1455	LRACSLPSMSALEKSMHLGRLPSRPPLPGSGGSQ SGAKMRMGPGRKRDFSPVPWSQYFESMEDVEV ENETGKDTFRVYKSGSEGPVLLLLHGGGHSALS WAVFTAAIISRVQCRIVALDLRSHGETKVKNPED LSAETMAKDVGNVVEAMYGDLPPPIMLIGHSMG GAIAVHTASSNLVPSLLGLCMIDVVEGTAMDAL NSMQNFLRGRPKTFKSLENAIEWSVKSGQIRNLE SARVSMVGQVKQCEGITSPEGSKSIVEGIIEEEEE DEEGSESISKRKKEDDMETKKDHPYTWRIELAKT EKYWDGWFRGLSNLFLSCPIPKLLLLAGVDRLD KDLTIGQMQGKFQMQVLPQCGHAVHEDAPDKV AEAVATFLIRHRFAEPIGGFQCVFPGC
3025	A	621	306	YHGGQRGRAGGSFRSVQGWGGQLRNPFRTSKSL SWKGLSSLLFPLYNLQMGRPRDRKELGRGHSPP HLEGPHMLPSGAARWRWLEAPVLVLEPLVLRPA AAPTP
3026	A	1533	454	AKVPQSTREEKRENGLEARSPAINLMGFNVEEM YEAHAWIQRILSLQNHHIIENNHILYLGRKEHDIL SQLQKTSSVSITEIISPGRTELEIEGARADLIEVVM NIEDMLCKVQEEMARKKERGLWRSLGQWTIQQ QKTQDEMKENIIFLKCPVPPTQELLDQKKQFEKC GLQVLKVEKIDNEVLMAAFQRKKKMMEEKLHR QPVSHRLFQQVPYQFCNVVCRVGFQRMYSTPCD PKYGAGIYFTKNLKNLAEKAKKISAADKLIYVFE AEVLTGFFCQGHPLNIVPPPLSPGAIDGHDSVVD NVSSPETFVIFSGMQAIPQYLWTCTQEYVQSQDY SSGPMRPFAQHPWRGFASGSPVD
3027	A .	179	703	PFHLGASSNTFRLQVQTQESKAQKEVKMGFIFSK SMNESMKNQKEFMLMNARLQLERQLIMQSEMR ERQMAMQIAWSREFLKYFGTFFGLAAISLTAGAI KKKKPAFLVPIVPLSFILTYQYDLGYGTLLERMK GEAEDILETEKSKLQLPRGMITFESIEKARKEQSR FFIDK
3028	A	876	1226	AVGKEPESSSTWVRDREGHIRSRRSMKMLWKLT DNIKYEDCEVSATPARSSVRSQAPSLTLPLLLLSL QPAAKRGWDKLSPAQRPSLGFARRTRGRSCRER TWMLPSLVSEFLHRD

SEQ ID NO:	Method	Predicted beginning nucleotide location corresponding to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion
3029	A	3	1731	FREGRFGSSCAVAAPLAGFQGLIECGYLAVDSPP SCWTPGGSNPAAPLPQALLPPRLPPTVLPFLGPGL SGELEMFTLPQKDFRAPTTCLGPTCMQDLGSSHG EDLEGECSRKLDQKLPELRGVGDPAMISSNTSYL SSRGRMIKWFWDSAEEGYRTYHMDEYDEDKNP SGIINLGTSENKLCFDLLSWRLSQRDMQRVEPSL LQYADWRGHLFLREEVAKFLSFYCKSPVPLRPE NVVVLNGGASLFSALATVLCEAGEAFLIPTPYYG AITQHVCLYGNIRLAYVYLDSEVTGLDTRPFQLT VEKLEMALREAHSEGVKVKGLILISPQNPLGDVY SPEELQEYLVFAKRHRLHVIVDEVYMLSVFEKSV GYRSVLSLERLPDPQRTHVMWATSKDFGMSGLR FGTLYTENQDVATAVASLCRYHGLSGLVQYQM AQLLRDRDWINQVYLPENHARLKAAHTYVSEEL RALGIPFLSRGAGFFIWVDLRKYLLKGTFEEEML LWRRFLDNKVLLSFGKAFECKEPGWFRFVFSDQ VHRLCLGMQRVQQVLAGKSQVAEDPRPSQSQEP SDQRR
3030	A .	1	584	PWLPWSDGRAARSSRKCPRSRFPVQVGKMAVST VFSTSSLMLALSRHSLLSPLLSVTSFRRFYRGDSP TDSQKDMIEIPLPPWQERTDESIETKRARLLYESR KRGMLENCILLSLFAKEHLQHMTEKQLNLYDRLI NEPSNDWDIYYWATEAKPAPEIFENEVMALLRD FAKNKNKEQRLRAPDLEYLFEKPR
3031	A		359	SLWPWILMDDSLMQISLQLLCVYTANFPNGCSSL CWSSCGQHPVQATHRGAVSNSLMLCILKLASQM PLENTTVQQMVFMLLSNLALSHDCKGVIQKSNF LQNFLSLALPKGGNKHLSNLTILWLKLLLNISSGE DGQQMILRLDGCLDLLTEMSKYKHKSSPLLPLLI FHNVCFSPANKPKILANEKVITVLAACLESENQN AQRIGAAALWALIYNYQKAKTALKSPSVKRRVD EAYSLAKKTFPNSEANPLNAYYLKCLENLVQLL NSS
3032	A		1242	GISGRPPRPAKRRMGKNPVRPPRALPPVPSQDDIP LSRPKKKKPRTKNTPASASLEGLAQTAGRRPSEG NEPSTKELKEHPEAPVQRRQKKTRLPLELETSST QKKSSSSSLLRNENGIDAEPAEEAVIQKPRRKTK KTQPAELQYANELGVEDEDIITDEQTTVEQQSVF TAPTGISQPVGKVFVEKSRRFQAADRSELIKTTEN IDVSMDVKPSWTTRDVALTVHRAFRMIGLFSHG FLAGCAVWNIVVIYVLAGDQLSNLSNLLQQYKT LAYPFQSLLYLLLALSTISAFDRIDFAKISVAIRNF LALDPTALASFLYFTALILSLSQQMTSDRIHLYTP SSVNGSLWEAGIEEQILQPWIVVNLVVALLVGLS WLFLSYRPGMDLSEELMFSSEVEEYPDKEKEIKA SS
3033	A .	3	1436	TATSGGIWLRKWRCHWPRPLPQSCVGTEGGLQ VRDTSSRIAKGGVDHTKMSLHGASGGHERSRDR RRSSDRSRDSSHERTESQLTPCIRNVTSPTRQHHV EREKDHSSSRPSSPRPQKASPNGSISSAGNSSRNS SQSSSDGSCKTAGEMVFVYENAKEGARNIRTSER VTLIVDNTRFVVDPSIFTAQPNTMLGRMFGSGRE HNFTRPNEKGEYEVAEGIGSTVFRAILDYYKTGII RCPDGISIPELREACDYLCISFEYSTIKCRDLSALM

HELSNDGARRQFEFYLEEMILPLMVASAQSGERE

SEQ ID NO:	Method	Predicted beginning nucleotide location corresponding to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion
				CHIVVLTDDDVVDWDEEYPPQMGEEYSQIIYSTK LYRFFKYIENRDVAKSVLKERGLKKIRLGIEGYP TYKEKVKKRPGGRPEVIYNYVQRPFIRMSWEKE EGKSRHVDFQCVKSKSITNLAAAAADIPQDQLV VMHPTPQVDELDILPIHPPSGNSDLDPDAQNPML
3034		3	1972	SSLAQHRSVAVLGWPAGWAAARARPAMQGGN SGVRKREEEGDGAGAVAAPPAIDFPAEGPDPEY DESDVPAEIQVLKEPLQQPTFPFAVANQLLLVSL LEHLSHVHEPNPLRSRQVFKLLCQTFIKMGLLSSF TCSDEFSSLRLHHNRAITHLMRSAKERVRQDPCE DISRIQKIRSREVALEAQTSRYLNEFEELAILGKG GYGRVYKVRNKLDGQYYAIKKILIKGATKTVCM KVLREVKVLAGLQHPNIVGYHTAWIEHVHVIQP RADRAAIELPSLEVLSDQEEDREQCGVKNDESSS SSIIFAEPTPEKEKRFGESDTENQNNKSVKYTTNL VIRESGELESTLELQENGLAGLSASSIVEQQLPLR RNSHLEESFTSTEESSEENVNFLGQTEAQYHLML HIQMQLCELSLWDWIVERNKRGREYVDESACPY VMANVATKIFQELVEGVFYIHNMGIVHRDLKPR NIFLHGPDQQVKIGDFGLACTDILQKNTDWTNR NGKRTPTHTSRVGTCLYASPEQLEGSEYDAKSD MYSLGVVLLELFQPFGTEMERAEVLTGLRTGQL PESLRKRCPVQAKYIQHLTRRNSSQRPSAIQLLQS ELFQNSGNVNLTLQMKIIEQEKEIAELKKQLNLL SQDKGVRDDGKDGGVG
3035	A	110	1172	KLSCPCSHGTRVTAVRGPRLKAGVQWHDLGSLQ PPPSGLKQSSHLSLSSSWDFRHAPTHPETYTCPK MIEMEQAEAQLAELDLLASMFPGENELIVNDQL AVAELKDCIEKKTMEGRSSKVYFTINMNLDVSD EKMAMFSLACILPFKYPAVLPEITVRSVLLSRSQQ TQLNTDLTAFLQKHCHGDVCILNATEWVREHAS GYVSRDTSSSPTTGSTVQSVDLIFTRLWIYSHHIY NKCKRKNILEWAKELSLSGFSMPGKPGVVCVEG PQSACEEFWARLRKLNWKRILIRHREDIPFDGTN DETERQRKFSIFEEKVFSVNGARGNHMDFGQLY QFLNTKGCGDVFQMFLWV
3036	A	1	2288	FRFAERRAAAAESDVSAKMAGRSMQAARCPTD ELSLTNCAVVNEKDFQSGQHVIVRTSPNHRYTFT LKTHPSVVPGSIAFSLPQRKWAGLSIGQEIEVSLY TFDKAKQCIGTMTIEIDFLQKKSIDSNPYDTDKM AAEFIQQFNNQAFSVGQQLVFSFNEKLFGLLVKD IEAMDPSILNGEPATGKRQKIEVGLVVGNSQVAF EKAENSSLNLIGKAKTKENRQSIINPDWNFEKMG IGGLDKEFSDIFRRAFASRVFPPEIVEQMGCKHVK GILLYGPPGCGKTLLARQIGKMLNAREPKVVNG PEILNKYVGESEANIRKLFADAEEEQRRLGANSG LHIIIFDEIDAICKQRGSMAGSTGVHDTVVNQLLS KIDGVEQLNNILVIGMTNRPDLIDEALLRPGRLEV KMEIGLPDEKGRLQILHIHTARMRGHQLLSADV DIKELAVETKNFSGAELEGLVRAAQSTAMNRHI KASTKVEVDMEKAESLQVTRGDFLASLENDIKP AFGTNQEDYASYIMNGIIKWGDPVTRVLDDGEL LVQQTKNSDRTPLVSVLLEGPPHSGKTALAAKIA EESNFPFIKICSPDKMIGFSETAKCQAMKKIFDDA YKSQLSCVVVDDIERLLDYVPIGPRFSNLVLQAL

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SEQ ID NO:	Method	Predicted beginning nucleotide location corresponding to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=-possible nucleotide insertion
				LVLLKKAPPQGRKLLIIGTTSRKDVLQEMEMLNA FSTTIHVPNIATGEQLLEALELLGNFKDKERTTIA QQVKGKKVWIGIKKLLMLIEMSLQMDPEYRVRK FLALLREEGASPLDFD
3037	A	I	1347	MLDTGSEHLNRILKALPALQSAGSEGQNGSAESL GEGGTRDSDRARRKLRGGNKEIPTFYPCLVVRSP VTASDLRGTQDFAAYHGLSLILEPLGACNRLSVC VPVHSPPGMRVSPRSPSLRTLVIDPAEPAGAQRL RFSGKERSGEAGSAVEGLAVAVSMGDGGAERD RGPARRAESGGGGGRCGDRSGAGDLRADGGGH SPTEVAGTSASSPAGSRESGADSDGQPGPGEADH CRRILVRDAKGTIREIVLPKGLDLDRPKRTRTFFT AEQLYRLEMEFQRCQYVVGRERTELARQLNLSE TQVKVWFQNRRTKQKKDQSRDLEKRASSSASEA FATSNILRLLEQGRLLSVPRAPSLLALTPSLPGLP ASHRGTSLGDPRNSSPRLNPLSSASASPPLPPPLP AVCFSSAPLLDLPAGYELGSSAFEPYSWLERKVG SASSCKKANT
3038	A	924	501	TELLPLCSRSGPKPQSGDPLLQLAQQARPRLSGE RLETAPSLLLSRMACVISGWALSRGARTWTWAT PTGPVHRAQPAIRSLSAEGALTRLKEEKWPGRYI LPNHLTPPFLYKHLGSVPPSHWRSPLISHSVNILA LNWR
3039	A	1263		ACGIRHEGALPGLTATPEAMLRFLPDLAFSFLLIL ALGQAVQFQEYVFLQFLGLDKAPSPQKFQPVPYI LKKIFQDREAAATTGVSRDLCYVKELGVRGNVL RFLPDQGFFLYPKKISQASSCLQKLLYFNLSAIKE REQLTLAQLGLDLGPNSYYNLGPELELALFLVQE PHVWGQTTPKPGKMFVLRSVPWPQGAVHFNLL DVAKDWNDNPRKNFGLFLEILVKEDRDSGVNFQ PEDTCARLRCSLHASLLVVTLNPDQCHPSRKRA AIPVPKLSCKNLCHRHQLFINFRDLGWHKWIIAP KGFMANYCHGECPFSLTISLNSSNYAFMQALMH AVDPEIPQAVCIPTKLSPISMLYQDNNDNVILRHY EDMVVDECGCG
3040	A .	15	849	ASRLPRGPGCGADMRPLLGLLLVFAGCTFALYL LSTRLPRGRRLGSTEEAGGRSLWFPSDLAELREL SEVLREYRKEHQAYVFLLFCGAYLYKQGFAIPGS SFLNVLAGALFGPWLGLLLCCVLTSVGATCCYL LSSIFGKQLVVSYFPDKVALLQRKVEENRNSLFF FLLFLRLFPMTPNWFLNLSAPILNIPIVQFFFSVLI GLIPYNFICVQTGSILSTLTSLDALFSWDTVFKLL AIAMVALIPGTLIKKFSQKHLQLNETSTANHIHSR KDT
3041	A	1015	175	GLKRRLCFAKVGDVLGCLSLPPSRSARVLEDISI LSCISVDSRIVRTKVPCSVTMSRPRKRLAGTSGSD KGLSGKRTKTENSGEALAKVEDSNPQKTSATKN CLKNLSSHWLMKSEPESRLEKGVDVKFSIEDLKA QPKQTTCWDGVRNYQARNFLRAMKLGEEAFFY HSNCKEPGIAGLMKIVKEAYPDHTQFEKNNPHY DPSSKEDNPKWSMVDVQFVRMMKRFIPLAELKS YHQAHKATGGPLKNMVLFTRQRLSIQPLTQEEF DFVLSLEEKEPS
3042	A	1015	175	GLKRRLCFAKVGDVLGCLSLPPSRSARVLEDISI LSCISVDSRIVRTKVPCSVTMSRPRKRLAGTSGSD

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SEQ ID NO:	Method	Predicted beginning nucleotide location corresponding to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion
				KGLSGKRTKTENSGEALAKVEDSNPQKTSATKN CLKNLSSHWLMKSEPESRLEKGVDVKFSIEDLKA QPKQTTCWDGVRNYQARNFLRAMKLGEEAFFY HSNCKEPGIAGLMKIVKEAYPDHTQFEKNNPHY DPSSKEDNPKWSMVDVQFVRMMKRFIPLAELKS YHQAHKATGGPLKNMVLFTRQRLSIQPLTQEEF DFVLSLEEKEPS
3043	A	153	1133	VGTAPAPGGRDRAPAMGSFQLEDFAAGWIGGA ASVIVGHPLDTVKTRLQAGVGYGNTLSCIRVVY RRESMFGFFKGMSFPLASIAVYNSVVFGVFSNTQ RFLSQHRCGEPEASPPRTLSDLLLASMVAGVVSV GLGGPVDLIKIRLQMQTQPFRDANLGLKSRAVAP AEQPAYQGPVHCITTIVRNEGLAGLYRGASAML LRDVPGYCLYFIPYVFLSEWITPEACTGPSPCAV WLAGGMAGAISWGTATPMDVVKSRLQADGVY LNKYKGVLDCISQSYQKEGLKVFFRGITVNAVR GFPMSAAMFLGYELSLQAIRGDHAVTSP
3044	A	41	1316	PPLGAGAGIHARSPHPARRLRLTAAGVGGRASG LLPTPWRRHHGPSGAAPYPAARLWQGPWRCRR PQPMAQRYDELPHYPGIADGPAALAGFPEAVPA APGPYGPHRPPQPLPPGLDSDGLKRDKDEIYGHP LFPLLALGFEKCELATCSPRDGAGAGLGTPRGGD VCSSDSFNEDNTAFAKQVCSERPFSSNPELDNLM IQAIQVLRFHLLELEKGKMPIDLVIEDRDGGCRE DFEDYPAPCPSLPDQNNIWIRDHEDSGSVHLGTP GPSSGGLASQSGDNSSDQGVGLDTSVASPSSGGE DEDLDQEPRRNKKRGIFPKVATNIMRAWLFQHL SHPYPSEEQKKQLAQDTGLTILQVNNWFINARRR IVQPMIDQSNRTGQGAAFSPEGQPIGGYTETEPH VAFRAPASVGMSLNSEGEWHYL
3045	A	3	967	VAHTQWHTCQRLSQLTHRSILKYLLIDTHACQV LILKHTHASLSLPSCQECFPSSIPSASHMVSHPHPP PSPRWGQTPEGLPAASPCGPGPRSCFSSILPTGDS WGMLACLCTVLWHLPAVPALNRTGDPGPGPSIQ KTYDLTRYLEHQLRSLAGTYLNYLGPPFNEPDFN PPRLGAETLPRATVDLEVWRSLNDKLRLTQNYE AYSHLLCYLRGLNRQAATAELRRSLAHFCTSLQ GLLGSIAGVMAALGYPLPQPLPGTEPTWTPGPAH SDFLQKMDDFWLLKELQTWLWRSAKDFNRLKK KMQPPAAAVTLHLGAHGF MYAYMYICTHICICAYRGIHIDVYLYMCIYIHIWI
3046	A	1185	1584	HTYLCVHIYVYVYICTHICMCIHTYVYVYTYMY VYTYICLCVYICLCVHIYLCVYIHMYMCTHICMC IHTYVHMCICVYIHMYTCVYVYTYTCVYMY
3047	A	811	132	SLDLLGPIGILQEGRDPGTQGPQEKEKQMPASPM NTDAHLDINFKEGLKKERSYTGQFEANVRDEER QCGCGVVPDSLLMKVLSQRLDQQDCIQKGWVL HGVPRDLDQAHLLNRLGYNPNREFFLNVPFDSI MERLTLRRIDPVTGERYHLMYKPPPTMEIQARLL QNPKDAEEQVKLKMDLFYRNSADLEQLYGSAIT LNGDQDPYTVFEYIESGIINPLPKKIP
3048	A	2	1166	RPRRGQGLVQEVQTENVTVAEGGVAEITCRLHQ YDGSIVVIQNPARQTLFFNGTRALKDERFQLEEFS PRRVRIRLSDARLEDEGGYFCQLYTEDTHHQIAT LTVLVAPENPVVEVREQAVEGGEVELSCLVPRSR

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				PAATLRWYRDRKELKGVSSSQENGKVWSVAST VRFRVDRKDDGGIIICEAQNQALPSGHSKQTQYV LDVQYSPTARIHASQAVVREGDTLVLTCAVTGN PRPNQIRWNRGNESLPERAEAVGETLTLPGLVSA DNGTYTCEASNKHGHARALYVLVVYGESRLRPT EGGGGAPDPGAVVEAQTSVPYAIVGGILALLVFL IICVLVGMVWCSVRQKGSYLTHEASGLDEQGEA REAFLNGSDGHKRKEEFFI
3049	A	3159	882	VGCTLRVGVMAAAGSRKRRLAELTVDEFLASGF DSESESESENSPQAETREAREAARSPDKPGGSPSA SRRKGRASEHKDQLSRLKDRDPEFYKFLQENDQ SLLNFSDSDSSEEEEGPFHSLPDVLEEASEEEDGA EEGEDGDRVPRGLKGKKNSVPVTVAMVERWKQ AAKQRLTPKLFHEVVQAFRAAVATTRGDQESAE ANKFQVTDSAAFNALVTFCIRDLIGCLQKLLFGK VAKDSSRMLQPSSSPLWGKLRVDIKAYLGSAIQL VSCLSETTVLAAVLRHISVLVPCFLTFPKQCRML LKRMVVVWSTGEESLRVLAFLVLSRVCRHKKDT FLGPVLKQMYITYVRNCKFTSPGALPFISFMQWT LTELLALEPGVAYQHAFLYIRQLAIHLRNAMTTR KKETYQSVYNWQYVHCLFLWCRVLSTAGPSEA LQPLVYPLAQVIIGCIKLIPTARFYPLRMHCIRALT LLSGSSGAFIPVLPFILEMFQQVDFNRKPGRMSSK PINFSVILKLSNVNLQEKAYRDGLVEQLYDLTLE YLHSQAHCIGFPELVLPVVLQLKSFLRECKVANY CRQVQQLLGKVQENSAYICSRRQRVSFGVSEQQ AVEAWEKLTREEGTPLTLYYSHWRKLRDREIQL EISGKERLEDLNFPEIKRKMADRKDEDRKQFKD LFDLNSSEEDDTEGFSERGILRPLSTRHGVEDDEE DEEEGEEDSSNSEDGDPDAEAGLAPGELQQLAQ GPEDELEDLQLSEDD
3050	A	870	182	HLDRYIKSPGSGSSTPAPPSHLLLYLLHPQSTRTM GCCGCSRGCGSGCGGCGSGCGGCGSG RGGCGSGCGGCGSSCGGCGSGCGGCGSG RGGCGSGCGCSSSCGGCGSRCYVPVCCCKPVC SWVPACSCTSCGSCGGSKGGCGSCGGSKGGCGS CGCSQSSCCKPCCCSSGCGSSCSQSSCCKPCCCSS GCGSSCCQSSCCKPYCCQSSCCKPCSCFSGCGSS CCQSSCYKPCCCQSSCCVPVCCQCKI
3051	A	175	4330	NIPRWNFQGKSFGVVLVHFSSEEVDMASDSPARS LDEIDLSALRDPAGIFELVELVGNGTYGQVYKGR HVKTGQLAAIKVMDVTGDEEEEIKQEINMLKKY SHHRNIATYYGAFIKKNPPGMDDQLWLVMEFCG AGSVTDLIKNTKGYTLKEEWIAYICREILRGLSHL HQHKVIHRDIKGQNVLLTENAEVKLVDFGVSAQ LDRTVGRRNTFIGTPYWMAPEVIACDENPDATY DFKSDLWSLGITAIEMAEGAPPLCDMHPMRALF LIPRNPAPRLKSKKWSKKFQSFIESCLVKNHSQRP ATEQLMKHPFIRDQPNERQVRIQLKDHIDRTKKK RGEKDETEYEYSGSEEEEEENDSGEPSSILNLPGE STLRRDFLRLQLANKERSEALRRQQLEQQREN EEHKRQLLAERQKRIEEQKEQRRRLEEQQRREKE LRKQQEREQRRHYEEQMRREEERRAEHEQEYI RRQLEEEQRQLEILQQQLLHEQALLLEYKRKQLE EQRQAERLQRQLKQERDYLVSLQHQRQEQRPVE KKPLYHYKEGMSPSEKPAWAKEVEERSRLNRQS

SEQ ID NO:	Method	Predicted beginning nucleotide location corresponding to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \possible nucleotide insertion
				SPAMPHKVANRISDPNLPPRSESFSISGVQPARTP PMLRPVDPQIPHLVAVKSQGPALTASQSVHEQPT KGLSGFQEALNVTSHRVEMPRQNSDPTSENPPLP TRIEKFDRSSWLRQEEDIPPKVPQRTTSISPALAR KNSPGNGSALGPRLGSQPIRASNPDLRRTEPILES PLQRTSSGSSSSSSSTPSSQPSSQGGSQPGSQAGSSE RTRVRANSKSEGSPVLPHEPAKVKPEESRDITRPS RPASYKKAIDEDLTALAKELRELRIEETNRPMKK VTDYSSSSEESESSEEEEEDGESETHDGTVAVSDI PRLIPTGAPGSNEQYNVGMVGTHGLETSHADSFS GSISREGTLMIRETSGEKKRSGHSDSNGFAGHINL PDLVQQSHSPAGTPTEGLGRVSTHSQEMDSGTE YGMGSSTKASFTPFVDPRVYQTSPTDEDEEDEES SAAALFTSELLRQEQAKLNEARKISVVNVNPTNI RPHSDTPEIRKYKKRFNSEILCAALWGVNLLVGT ENGLMLLDRSGQGKVYNLINRRRFQQMDVLEG LNVLVTISGKKNKLRVYYLSWLRNRILHNDPEV EKKQGWITVGDLEGCIHYKVVKYERIKFLVIALK NAVEIYAWAPKPYHKFMAFKSFADLQHKPLLVD LTVEEGQRLKVIFGSHTGFHVIDVDSGNSYDIYIP SHIQGNITPHAIVILPKTDGMEMLVCYEDEGVYV NTYGRITKDVVLQWGEMPTSVAYIHSNQIMGW GEKAIEIRSVETGHLDGVFMHKRAQRLKFLCERN DKVFFASVRSGGSSQVFFMTLNRNSMMNW
3052	Α	1	615	MGQVECGGQKLGNQLEDDSEPAEGKVYSSDEE KLEASAGDPAGSEQEEEGSGGDSEDDGFLDSSA GGPGALLGPKPKLKGSLGTGAEEGAPVTAGVTA PGGKSRRRRTAFTSEQLLELEKEFHCKKYLSLTE RSQIAHALKLSEVQVKIWFQNRRAKWKRIKAGN VSSRSGEPVRNPKIVVPIPVHVNRFAVRSQHQQM EQGARP
3053	A .	203	2167	FGVRVPSNTQCLVPSFHCMQTSEWDSECLTSLQP LPLPTPPAANEAHLQTAAISLWTVVAAVQAIERK VEIHSRRLLHLEGRTGTAEKKLASCEKTVTELGN QLEGKGAVLGTLLQEYGLLQRRLENLENLLRNR NFWILRLPPGIKGDIPKVPVAFDDVSIYFSTPEWE KLEEWQKELYKNIMKGNYESLISMDYAINQPDV LSQIQPEGEHNTEDQAGPEESEIPTDPSEEPGISTS DILSWIKQEEEPQVGAPPESKESDVYKSTYADEE LVIKAEGLARSSLCPEVPVPFSSPPAAAKDAFSDV AFKSQQSTSMTPFGRPATDLPEASEGQVTFTQLG SYPLPPPVGEQVFSCHHCGKNLSQDMLLTHQCS HATEHPLPCAQCPKHFTPQADLSSTSQDHASETP PTCPHCARTFTHPSRLTYHLRVHNSTERPFPCPDC PKRFADQARLTSHRRAHASERPFRCAQCGRSFSL KISLLLHQRGHAQERPFSCPQCGIDFNGHSALIRH QMIHTGERPYPCTDCSKSFMRKEHLLNHRRLHT GERPFSCPHCGKSFIRKHHLMKHQRIHTGERPYP CSYCGRSFRYKQTLKDHLRSGHNGGCGGDSDPS GQPPNPPGPLITGLETSGLGVNTEGLETNQWYGE GSGGGVL
3054	A	3	2212	SCGHKSAYGSYTGLQLFWEDGQELLQHQQLQD LRLCVHLRPQSEKVELSLWTLFVVGKGEPSAVR EKLGKAGFAAASGPGGRPGAERASTVLNILHLT AESRWEPNACNRVSSSPAGVGPLDLPVGPLLYFF

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SEQ ID NO:	Method	Predicted beginning nucleotide location corresponding to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion
				APWARASFLCHAFQRPLTGIGLNTVRFTSEFPLH SKDPTAHKLLFTGNYLCKLHPRPRHAPQGSLSDF CHGTEGKDLPSEHNVSVEGVAQDRSPEATLCPQ KTCPCDICGLRLKDILHLAEHQTTHPRQKPFVCE AYVKGSEFSANLPRKQVQQNVHNPIRTEEGQAS PVKTCRDHTSDQLSTCREGGKDFVATAGFLQCE VTPSDGEPHEATEGVVDFHIALRHNKCCESGDAF NNKSTLVQHQRIHSRERPYECSKCGIFFTYAADL TQHQKVHNRGKPYECCECGKFFSQHSSLVKHRR
				VHTGESPHVCGDCGKFFSRSSNLIQHKRVHTGEK PYECSDCGKFFSQRSNLIHHKRVHTGRSAHECSE CGKSFNCNSSLIKHWRVHTGERPYKCNECGKFFS HIASLIQHQIVHTGERPHGCGECGKAFIRSSDLMK HQRVHTGERPYECNECGKLFSQSSSLNSHRRLHT GERPYQCSECGKFFNQSSSLNNHRRLHTGERPYE CSECGKTFRQRSNLRQHLKVHKPDRPYECSECG KAFNQRPTLIRHQKIHIRERSMENVLLPCSQHTPE ISSENRPYQGAVNYKLKLVHPSTHPGEVP
3055	A	268	2954	ARRSSSSQGSAAPTPCQVVEASRDQLVAGPSGK MGNREMEELIPLVNRLQDAFSALGQSCLLELPQI AVVGGQSAGKSSVLENFVGRDFLPRGSGIVTRRP LVLQLVTSKAEYAEFLHCKGKKFTDFDEVRLEIE AETDRVTGMNKGISSIPINLRVYSPHVLNLTLIDL PGITKVPVGDQPPDIEYQIRMIMQFITRENCLILA VTPANTDLANSDALKLAKEVDPQGLRTIGVITKL DLMDEGTDARDVLENKLLPLRRGYVGVVNRSQ KDIDGKKDIKAAMLAERKFFLSHPAYRHIADRM GTPHLQKVLNQQLTNHIRDTLPNFRNKLQGQLLS IEHEVEAYKNFKPEDPTRKTKALLQMVQQFAVD FEKRIEGSGDQVDTLELSGGAKINRIFHERFPFEIV KMEFNEKELRREISYAIKNIHGIRTGLFTPDMAFE AIVKKQIVKLKGPSLKSVDLVIQELINTVKKCTK KLANFPRLCEETERIVANHIREREGKTKDQVLLLI DIQVSYINTNHEDFIGFANAQQRSSQVHKKTTVG NQVIRKGWLTISNIGIMKGGSKGYWFVLTAESLS WYKDDEEKEKKYMLPLDNLKVRDVEKSFMSSK HIFALFNTEQRNVYKDYRFLELACDSQEDVDSW KASLLRAGVYPDKSVGNNKAENDENGQAENFS MDPQLERQVETIRNLVDSYMSIINKCIRDLIPKTI MHLMINNVKDFINSELLAQLYSSEDQNTLMEES AEQAQRRDEMLRMYQALKEALGIIGDIGTATVS TPAPPPVDDSWIQHSRRSPPPSPTTQRRPTLSAPL ARPTSGRGPAPAIPSPGPHSGAPPVPFRPGPLPPFP SSSDSFGAPPQVPSRPTRAPPSVPSRRPPPSPTRPTI IRPLESSLLD
3056	A	1674	1839	VVRVTCCPPARSTTERTNAYDEEDCVEMVASGG WNDVACHTTMYFMCEFDKKNM
3057	A	1674	1839	VVRVTCCPPARSTTERTNAYDEEDCVEMVASGG WNDVACHTTMYFMCEFDKKNM
3058	A	3363	2525	FLVKLILIILCRCLHSLSRSVQQLRTSFQDHAVWK PLMKVLQNAPDEILVVASSMLCNLLLEFSPSKEPI LESGAVELLCGLTQSENPALRVNGIWALMNMAF QAEQKIKADILRSLSTEQLFRLLSDSDLNVLMKT LGLLRNLLSTRPHIDKIMSTHGKQIMQAVTLILEG EHNIEVKEQTLCILANIADGTTAKDLIMTNDDILQ

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SEQ ID NO:	Method	Predicted beginning nucleotide location corresponding to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion
				KIKYYMGHSHVKLQLAAMFCISNLIWNEEEGSQ ERQDKLRDMGIVDILHKLSQSPDSNLCDKAKMA LQQYLA
3059	A	679	167	SSWPSLSSQMHFPSFHLHVAAHYGRDSFVRLLLE FKAEVDPLSDKGTTPLQLAIIRERSSCVKILLDHN ANIDIQNGFLLRYAVIKSNHSYCRMFLQRGADTN LGRLEDGQTPLHLSALRDDVLCARMLYNYGAD TNTRNYEGQTPLAVSISISGSSRPCLDFLQEVTSM
3060	A	30	234	PPLQLDMDPNCYCADGDSCTCAGSCKCKECKCT SCKKSCCSCCPAGCAKCAQGCICKGATDKCSCC A
3061	A	428	720	VRRDVRQQATWAMASDLDFSPPEVPEPTFLENL LRYGLFLGAIFQLICVLAIIVPIPKSHEAEAEPSEPR SAEVTRKPKAAVPSVNKRPKKETKKKR
3062	A	1589	276	WKQKYEPLGLDAAGIEEAITAVGSFILKANELLQ VIDSSMKNFKAFFRWLYVAMLRMTEDHVLPELN KMTQKDITFVAEFLTEHFNEAPDLYNRKGKYFN VERVGQYLKDEDDDLVSPPNTEGNQWYDFLQN SSHLKESPLLFPYYPRKSLHFVKRRMENIIDQCLQ KPADVIGKSMNQAICIPLYRDTRSEDSTRRLFKFP FLWNNKTSNLHYLLFTILEDSLYKMCILRRHTDIS QSVSNGLIAIKFGSFTYATTEKVRRSIYSCLDAQF YDDETVTVVLKDTVGREGRDRLLVQLPLSLVYN SEDSAEYQFTGTYSTRLDEQCSAIPTRTMHFEKH
				WRLLESMKAQYVAGNGFRKVSCVLSSNLRHVR VFEMDIDDEWELDESSDEEEEASNKPVKIKEEVL SESEAENQQAGAAALAPEIVIKVEKLDPELDS
3063	A	50	849	DKMPSIFAYQSSEVDWCESNFQYSELVAEFYNTF SNIPFFIFGPLMMLLMHPYAQKRSRYIYVVWVLF MIIGLFSMYFHMTLSFLGQLLDEIAILWLLGSGYS IWMPRCYFPSFLGGNRSQFIRLVFITTVVSTLLSFL RPTVNAYALNSIALHILYIVCQEYRKTSNKELRH LIEVSVVLWAVALTSWISDRLLCSFWQRIHFFYL HSIWHVLISITFPYGMVTMALVDANYEMPGETL KVRYWPRDSWPVGLPYVEIRGDDKDC
3064	A	1523	925	AATMADGQMPFSCHYPSRLRRDPFRDSPLSSRLL DDGFGMDPFPDDLTASWPDWALPRLSSAWPGTL RSGMVPRGPTATARFGVPAEGRTPPPFPGEPWK VCVNVHSFKPEELMVKTKDGYVEVSGKHEEKQ QEGGIVSKNFTKKIQLPAEVDPVTVFASLSPEGLL IIEAPQVPPYSTFGESSFNNELPQDSQEVTCT
3065	A	230	2929	LSTSLTGSHLFSLGNHSTRENLNAGNFNFPSEGH LVRSTGPGGSFAKHMVAQCVSPKGPLACSRTYF FGATHVPYLGGDSKLPKKTEQIRLLSQIYAAVIE AVLAGIACYAKTSSLTKAKEVAEQTLGSGLDSFE LIPFKAALRSKMTFHIHAVNNQGRIVPLDSEDSLS FVKTACMAVYDIPDLLGGNGCLGSVVFSESFLTS QILVKEKDGTVTTETSSVVLTAAVPRFCSWLVED NEVKLSEKTHQAVRGDESFLGTYLTGGEGAYLY SSNLQSWPEEGNVHFFSSGLLFSHCRHGSIIISKD HMNSISFYDGDSTSTVAALLIDFKSSLLPHLPVHF HGSSNFLMIALFPKSKIYQAFYSEVFSLWKQQDN SGISLKVIQEDGLSVEQKRLHSSAQKLFSALSQPA GEKRSSLKLLSAKLPELDWFLQHFAISSISQEPVM RTHLPVLLQQAEINTTHRIESDKVIISIVTGLPGCH

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				ASELCAFLVTLHKECGRWMVYRQIMDSSECFHA AHFQRYLSSALEAQQNRSARQSAYIRKKTRLLV VLQGYTDVIDVVQALQTHPDSNVKASFTIGAITA CVEPMSCYMEHRFLFPKCLDQCSQGLVSNVVFT SHTTEQRHPLLVQLQSLIRAANPAAAFILAENGIV TRNEDIELILSENSFSSPEMLRSRYLMYPGWYEG KLNAGSVYPLMVQICVWFGRPLEKTRFVAKCKA IQSSIKPSPFSGNIYHILGKVKFSDSERTMEVCYNT LANSLSIMPVLEGPTPPPDSKSVSQDSSGQQECYL VFIGCSLKEDSIKDWLRQSAKQKPQRKALKTRG MLTQQEIRSIHVKRHLEPLPAGYFYNGTQFVNFF GDKTDFHPLMDQFMNDYVEEANREIEKYNQELE QQEYHDLFELKP
3066	A	130	588	LAPLRCQPGTRTQPRSHPAANDPSAAMSAAGAR GLRATYHRLLDKVELMLPEKLRPLYNHPAGPRT VFFWAPIMKWGLVCAGLADMARPAEKLSTAQS AVLMATGFIWSRYSLVIIPKNWSLFAVNFFVGAA GASQLFRIWRYNQELKAKAHK
3067	A	2	1016	EFARRVFIAAREMSLLRSLRVFLVARTGSYPAG SLLRQSPQPRHTFYAGPRLSASASSKELLMKLRR KTGYSFVNCKKALETCGGDLKQAEIWLHKEAQ KEGWSKAAKLQGRKTKEGLIGLLQEGNTTVLVE VNCETDFVSRNLKFQLLVQQVALGTMMHCQTL KDQPSAYSKGFLNSSELSGLPAGPDREGSLKDQL ALAIGKLGENMILKRAAWVKVPSGFYVGSYVHG AMQSPSLHKLVLGKYGALVICETSEQKTNLEDV GRRLGQHVVGMAPLSVGSLDDEPGGEAETKML SQPYLLDPSITLGQYVQPQGVSVVDFVRFECGEG EEAAETE
3068		3	1679	NSRVWGPWTEPSAGSLRPMARKQNRNSKELGL VPLTDDTSHAGPPGPGRALLECDHLRSGVPGGR RRKDWSCSLLVASLAGAFGSSFLYGYNLSVVNA PTPYIKAFYNESWERRHGRPIDPDTLTLLWSVTV SIFAIGGLVGTLIVKMIGKVLGRKHTLLANNGFAI SAALLMACSLQAGAFEMLIVGRFIMGIDGGVALS VLPMYLSEISPKEIRGSLGQVTAIFICIGVFTGQLL GLPELLGKESTWPYLFGVIVVPAVVQLLSLPFLP DSPRYLLLEKHNEARAVKAFQTFLGKADVSQEV EEVLAESRVQRSIRLVSVLELLRAPYVRWQVVT VIVTMACYQLCGLNAIWFYTNSIFGKAGIPPAKIP YVTLSTGGIETLAAVFSGLVIEHLGRRPLLIGGFG LMGLFFGTLTITLTLQDHAPWVPYLSIVGILAIIAS FCSGPGGIPFILTGEFFQQSQRPAAFIIAGTVNWLS NFAVGLLFPFIQKSLDTYCFLVFATICITGAIYLYF VLPETKNRTYAEISQAFSKRNKAYPPEEKIDSAV TDGKINGRP
3069	A .	861	300	AAGAVVSAMPKAKGKTRQKFGYSVNRKRLNR NARRKAAPRIECSHIRHAWDHAKSVRQNLAEMG LAVDPNRAVPLRKRKVKAMEVDIEERPKELVRK PYVLNDLEAEASLPEKKGNTLSRDLIDYVRYMV ENHGEDYKAMARDEKNYYQDTPKQIRSKINVY KRFYPAEWQDFLDSLQKRKMEVE
3070	A	325	2019	LAEPEVATDSGQQADLPAEGGDPRAEASCSVLH SKPHAMADSRDPASDQMQHWKEQRAAQKADV LTTGAGNPVGDKLNVITVGPRGPLLVQDVVFTD

SEQ ID	Method	Predicted	Predicted end	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine,
NO:		beginning	nucleotide	I=Isoleucine, K=Lysine, L=Leucine, M=Methionine,
ł		nucleotide location	location corresponding	N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine,
		corresponding	to last amino	T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine,
		to first amino	acid residue of	X=Unknown, *=Stop codon, /=possible nucleotide deletion,
		acid residue of	peptide	>=possible nucleotide insertion
Į.		peptide	sequence	
		sequence		
				EMAHFDRERIPERVVHAKGAGAFGYFEVTHDIT
	İ			KYSKAKVFEHIGKKTPIAVRFSTVAGESGSADTV
				RDPRGFAVKFYTEDGNWDLVGNNTPIFFIRDPILF
				PSFIHSQKRNPQTHLKDPDMVWDFWSLRPESLH
1			1	QVSFLFSDRGIPDGHRHMNGYGSHTFKLVNANG
				EAVYCKFHYKTDQGIKNLSVEDAARLSQEDPDY
				GIRDLFNAIATGKYPSWTFYIQVMTFNQAETFPF
				NPFDLTKVWPHKDYPLIPVGKLVLNRNPVNYFA
	ŀ			
	İ		ĺ	EVEQIAFDPSNMPPGIEASPDKMLQGRLFAYPDT
			<u> </u>	HRHRLGPNYLHIPVNCPYRARVANYQRDGPMC
				MQDNQGGAPNYYPNSFGAPEQQPSALEHSIQYS
	1			GEVRRFNTANDDNVTQVRAFYVNVLNEEQRKR
į	l			LCENIAGHLKDAQIFIQKKAVKNFTEVHPDYGSH
			i	IQALLDKYNAEKPKNAIHTFVQSGSHLAAREKA
				NL
3071	A	1	1187	SLGWLERPPALSRAAGDGARRLSGSRRGDVWLT
3071	1 🚡	*	1107	SSAAGLLRSVAGGSWCGGQLRARGGSGRCVAR
	1			AMTGNAGEWCLMESDPGVFTELIKGFGCRGAQ
ł				
	ļ			VEEIWSLEPENFEKLKPVHGLIFLFKWQPGEEPA
	l			GSVVQDSRLDTIFFAKQVINNACATQAIVSVLLN
	1	<u> </u>		CTHQDVHLGETLSEFKEFSQSFDAAMKGLALSN
				SDVIRQVHNSFARQQMFEFDTKTSAKEEDAFHF
1				VSYVPVNGRLYELDGLREGPIDLGACNQDDWIS
	ŀ	1	Ì	AVRPVIEKRIQKYSEGEIRFNLMAIVSDRKMIYEQ
ł		ľ		KIAELQRQLAEEEPMDTDQGNSMLSAIQSEVAK
	l			NQMLIEEEVQKLKRYKIENIRRKHNYLPFIMELL
			'	KTLAEHQQLIPLVEKAKEKQNAKKAQETK
3072	Α	103	2775	RLRTLAPPGLLLGPPLVPDSRRRHQASLTPLHISG
3072	Λ.	103	2775	SPOLVGRGDRKLRTEVLVPPAALPAETRQRRSER
				LPRRTCPRGGAPGPGRSRLPRSLPPPSAIPGLRSPV
1				WAAGLGGGGRREPSRGKGGAALRARHRSTMAE
	l			LGAGGDGHRGGDGAVRSETAPDSYKVQDKKNA
Ì				SSRPASAISGQNNNHSGNKPDPPPVLRVDDRQRL
				ARERREEREKQLAAREIVWLEREERARQHYEKH
		,		LEERKKRLEEQRQKEERRRAAVEEKRRQRLEED
				KERHEAVVRRTMERSQKPKQKHNRWSWGGSLH
]				GSPSIHSADPDRRSVSTMNLSKYVDPVISKRLSSS
				SATLLNSPDRARRLQLSPWESSVVNRLLTPTHSF
				LARSKSTAALSGEAVIPICPRSASCSPIIMPYKAAH
				SRNSMDRPKLFVTPPEGSSRRRIIHGTASYKKERE
				RENVLFLTSGTRRAVSPSNPKARQPARSRLWLPS
				KSLPHLPGTPRPTSSLPPGSVKAAPAQVRPPSPGN
	1			IRPVKREVKVEPEKKDPEKEPQKVANEPSLKGRA
				PLVKVEEATVEERTPAEPEVGPAAPAMAPAPAS
			1	APAPASAPAPAPVPTPAMVSAPSSTVNASASVKT
1				SAGTTDPEEATRLLAEKRRLAREQREKEERERRE
				QEELERQKREELAQRVAEERTTRREEESRRLEAE
				QAREKEEQLQRQAEERALREWEEAERAQRQKEE
ſ				EARVREEAERVRQEREKHFQREEQERLERKKRL
				EEIMKRTRRTEATDKKTSDQRNGDIAKGALTGG
				TEVSALPCTTNAPGNGKPVGSPHVVTSHQSKVT
		1		VESTPDLEKQPNENGVSVQNENFEEIINLPIGSKP
	1			
]			SRLDVTNSESPEIPLNPILAFDDEGTLGPLPQVDG
	ļ			VQTQQTAEVI
3073	Α	67	2415	PPRVCRDHVCLICWDPIAGTGGSRSTMPALPLDQ

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SEQ ID NO:	Method	Predicted beginning nucleotide location corresponding to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion
				LQITHKDPKTGKLRTSPALHPEQKADRYFVLYKP PPKDNIPALVEEYLERATFVANDLDWLLALPHD KFWCQVIFDETLQKCLDSYLRYVPRKFDEGVAS APEVVDMQKRLHRSVFLTFLRMSTHKESKDHFIS PSAFGEILYNNFLFDIPKILDLCVLFGKGNSPLLQ KMIGNIFTQQPSYYSDLDETLPTILQVFSNILQHC GLQGDGANTTPQKLEERGRLTPSDMPLLELKDIV LYLCDTCTTLWAFLDIFPLACQTFQKHDFCYRLA SFYEAAIPEMESAIKKRRLEDSKLLGDLWQRLSH SRKKLMEIFHIILNQICLLPILESSCDNIQGFIEEFL QIFSSLLQEKRFLRDYDALFPVAEDISLLQQASSV LDETRTAYILQAVESAWEGVDRRKATDAKDPSV IEEPNGEPNGVTVTAEAVSQASSHPENSEEEECM GAAAAVGPAMCGVELDSLISQVKDLLPDLGEGFI LACLEYYHYDPEQVINNILEERLAPTLSQLDRNL DREMKPDPTPLLTSRHNVFQNDEFDVFSRDSVDL SRVHKGKSTRKEENTRSLLNDKRAVAAQRQRYE QYSVVVEEVPLQPGESLPYHSVYYEDEYDDTYD GNQVGANDADSDDELISRRPFTIPQVLRTKVPRE GQEEDDDDEEDDADEEAPKPDHFVQDPAVLREK AEARMAFLAKKGYRHDSSTAVAGSPRGHGQS RETTQERRKKEANKATRANHNRRTMADRKRSK GMIPS
3074	Α .	3	251	GEARSPPPAAALLDMDPETCPCPSGGSCTCADSC KCEGCKCTSCKKSCCSCCPAECEKCAKDCVCKG GEAAEAEKCSCCQ
3075	A	255	982	SQFSLSQVLVDSAEEGSLAAAAELAAQKREQRL RKFRELHLMRNEARKLNHQEVVEEDKRLKLPAN WEAKKARLEWELKEEEKKKECAARGEDYEKVK LLEISAEDAERWERKKKRKNPDLGFSDYAAAQL RQYHRLTKQIKPDMETYERLREKHGEEFFPTSNS LLHGTHVPSTEEIDRMVIDLEKQIEKRDKYSRRR PYNDDADIDYINERNAKFNKKAERFYGKYTAEI KQNLERGTAV
3076		255	982	SQFSLSQVLVDSAEEGSLAAAAELAAQKREQRL RKFRELHLMRNEARKLNHQEVVEEDKRLKLPAN WEAKKARLEWELKEEEKKKECAARGEDYEKVK LLEISAEDAERWERKKKRKNPDLGFSDYAAAQL RQYHRLTKQIKPDMETYERLREKHGEEFFPTSNS LLHGTHVPSTEEIDRMVIDLEKQIEKRDKYSRRR PYNDDADIDYINERNAKFNKKAERFYGKYTAEI KQNLERGTAV
3077	A	1		FRLRPRRACAQLLWHPAAGMASWAKGRSYLAP GLLQGQVAIYTGGATGIGKAIVKELLELGSNVVI ASRKLERLKSAADELQANLPPTKQARVIPIQCNIR NEEEVNNLVKSTLDTFGKINFLVNNGGGQFLSPA EHISSKGWHAVLETNLTGTFYMCKAVYSSWMK KHGGSIVNIIVPTKAGFPLAVHSGAARAGVYNLT KSLAFEWACSGIRINCVAPGVIYSQTAVENYGSW GQSFFEGSFQKIPAKRIGVPEEVSSVVCFLLSPAA SFITGQSVDVDGGRSLYTHSYEVPDHDNWPKGA GDLSVVKKMKETFKEKAKL
3078	A	2	3508	FVRESGKAPVTFDDITVYLLQEEWVLLSQQQKEL CGSNKLVAPLGPTVANPELFRKFGRGPEPWLGS VQGQRSLLEHHPGKKQMGYMGEMEVQGPTRES

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Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid. SEQ ID Method Predicted end Predicted NO: beginning nucleotide E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, nucleotide location corresponding N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, location T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, to last amino corresponding X=Unknown, *=Stop codon, /=possible nucleotide deletion, to first amino acid residue of acid residue of peptide \=possible nucleotide insertion peptide sequence sequence GOSLPPOKKAYLSHLSTGSGHIEGDWAGRNRKL LKPRSIQKSWFVQFPWLIMNEEQTALFCSACREY PSIRDKRSRLIEGYTGPFKVETLKYHAKSKAHMF CVNALAARDPIWAARFRSIRDPPGDVLASPEPLF TADCPIFYPPGPLGGFDSMAELLPSSRAELEDPGG DGAIPAMYLDCISDLRQKEITDGIHSSSDINILYN DAVESCIQDPSAEGLSEEVPVVFEELPVVFEDVA VYFTREEWGMLDKROKELYRDVMRMNYELLAS LGPAAAKPDLISKLERRAAPWIKDPNGPKWGKG RPPGNKKMVAVREADTQASAADSALLPGSPVEA RASCCSSSICEEGDGPRRIKRTYRPRSIQRSWFGQ FPWLVIDPKETKLFCSACIERPNLHDKSSRLVRG YTGPFKVETLKYHEVSKAHRLCVNTVEIKEDTPH TALVPEISSDLMANMEHFFNAAYSIAYHSRPLND FEKILQLLQSTGTVILGKYRNRTACTQFIKYISETL KREILEDVRNSPCVSVLLDSSTDASEQACVGIYIR YFKQMEVKESYITLAPLYSETADGYFETIVSALD ELDIPFRKPGWVVGLGTDGSAMLSCRGGLVEKF **QEVIPOLLPVHCVAHRLHLAVVDACGSIDLVKK** CDRHIRTVFKFYQSSNKRLNELQEGAAPLEQEIIR LKDLNAVRWVASRRRTLHALLVSWPALARHLQ RVAEAGGOIGHRAKGMLKLMRGFHFVKFCHFL LDFLSIYRPLSEVCQKEIVLITEVNATLGRAYVAL **ESLRHQAGPKEEEFNASFKDGRLHGICLDKLEVA EQRFQADRERTVLTGIEYLQQRFDADRPPQLKN** MEVFDTMAWPSGIELASFGNDDILNLARYFECSL PTGYSEEALLEEWLGLKTIAQHLPFSMLCKNALA OHCRFPLLSKLMAVVVCVPISTSCCERGFKAMN RIRTDERTKLSNEVLNMLMMTAVNGVAVTEYD POPAIOHWYLTSSGRRFSHVYTCAQVPARSPASA RLRKEEMGALYVEEPRTQKPPILPSREAAEVLKD CIMEPPERLLYPHTSOEAPGMS 3079 1513 **FSPLEPRLCSLGGWGALOAGEPCOPSRAGCGRE** A 343 GATMGCTLSAEERAALERSKAIEKNLKEDGISAA KDVKLLLLGAGESGKSTIVKQMKIIHEDGFSGED VKQYKPVVYSNTIQSLAAIVRAMDTLGIEYGDK **ERKADAKMVCDVVSRMEDTEPFSAELLSAMMR** LWGDSGIQECFNRSREYQLNDSAKYYLDSLDRIG AADYQPTEQDILRTRVKTTGIVETHFTFKNLHFR LFDVGGQRSERKKWIHCFEDVTAIIFCVALSGYD OVLHEDETTNRMHESLKLFDSICNNKWFTDTSII LFLNKKDIFEEKIKKSPLTICFPEYTGPSAFTEAVA YIQAQYESKNKSAHKEIYSHVTCATDTNNIQFVF DAVTDVIIAKNLRGCGLY 3080 41 997 EARTARELTDGVTDGLTMADOPKPISPLKNLLA Α GGFGGVCLVFVGHPLDTVKVRLQTQPPSLPGQPP MYSGTFDCFRKTLFREGITGLYRGMAAPIIGVTP MFAVCFFGFGLGKKLQQKHPEDVLSYPQLFAAG MLSGVFTTGIMTPGERIKCLLQIQASSGESKYTGT LDCAKKLYQEFGIRGIYKGTVLTLMRDVPASGM YFMTYEWLKNIFTPEGKRVSELSAPRILVAGGIA **GIFNWAVAIPPDVLKSRFQTAPPGKYPNGFRDVL** RELIRDEGVTSLYKGFNAVMIRAFPANAACFLGF **EVAMKFLNWATPNL**

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IMADMEDLFGSDADSEAERKDSDSGSDSDSDQE

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SEQ ID NO:	Method	Predicted beginning	Predicted end nucleotide	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine,
		nucleotide	location	I=Isoleucine, K=Lysine, L=Leucine, M=Methionine,
1		location	corresponding	N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine,
		corresponding	to last amino	T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine,
		to first amino acid residue of	acid residue of peptide	X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion
i		peptide	sequence	- Possible nucleotitue insertion
		sequence		
				NAASGSNASGSESDQDERGDSGQPSNKELFGDD
}	ļ		1	SEDEGASHHSGSDNHSERSDNRSEASERSDHEDN
				DPSDVDQHSGSEAPNDDEDEGHRSDGGSHHSEA
				EGSEKAHSDDEKWGREDKSDQSDDEKIQNSDDE
				ERAQGSDEDKLQNSDDDEKMQNTDDEERPQLS
				DDERQQLSEEEKANSDDERPVASDNDDEKQNSD
			}	DEEQPQLSDEEKMQNSDDERPQASDEEHRHSDD
				EEEQDHKSESARGSDSEDEVLRMKRKNAIASDSE
				ADSDTEVPKDNSGTMDLFGGADDISSGSDGEDK
			J	PPTPGQPVDENGLPQDQQEEEPIPETRIEVEIPKV
İ				NTDLGNDLYFVKLPNFLSVEPRPFDPQYYEDEFE
		Į.		DEEMLDEEGRTRLKLKVENTIRWRIRRDEEGNEI
				KESNARIVKWSDGSMSLHLGNEVFDVYKAPLQG
				DHNHLFIRQGTGLQGQAVFKTKLTFRPHSTDSAT
	1]	HRKMTLSLADRCSKTQKIRILPMAGRDPECQRTE
			1	MIKKEEERLRASIRRESQQRRMREKQHQRGLSAS
				YLEPDRYDEEEEGEESISLAAIKNRYKGGIREERA
				RIYSSDSDEGSEEDKAQRLLKAKKLTSDEVRPNL
	ĺ			FNSRGLSCTQEPTALNEELTDQAGTN
3082	A	3	921	VEFCLPASADSSSLVAASLAGVRKMATNFLAHE
		ļ	•	KIWFDKFKYDDAERRFYEQMNGPVAGASRQEN
		j		GASVILRDIARARENIQKSLAGSSGPGASSGTSGD
				HGELVVRIASLEVENQSLRGVVQELQQAISKLEA
				RLNVLEKSSPGHRATAPQTQHVSPMRQVEPPAK
				KPATPAEDDEDDDIDLFGSDNEEEDKEAAQLREE
				RLRQYAEKKAKKPALVAKSSILLDVKPWDDETD
	0			MAQLEACVRSIQLDGLVWGASKLVPVGYGIRKL
				QIQCVVEDDKVGTDLLEEEITKFEEHVQSVDIAA
				FNKI
3083	A	3	921	VEFCLPASADSSSLVAASLAGVRKMATNFLAHE
}				KIWFDKFKYDDAERRFYEQMNGPVAGASRQEN
				GASVILRDIARARENIQKSLAGSSGPGASSGTSGD
				HGELVVRIASLEVENQSLRGVVQELQQAISKLEA
1				RLNVLEKSSPGHRATAPQTQHVSPMRQVEPPAK
			İ	KPATPAEDDEDDDIDLFGSDNEEEDKEAAQLREE
				RLRQYAEKKAKKPALVAKSSILLDVKPWDDETD
		,		MAQLEACVRSIQLDGLVWGASKLVPVGYGIRKL
ĺ		ĺ		QIQCVVEDDKVGTDLLEEEITKFEEHVQSVDIAA
2004	<u> </u>	120	4050	FNKI VENWINVEMA A ETOTI NECDENII DAI SECCEITE
3084	A	128	4050	KSIVKIRKRMAAETQTLNFGPEWLRALSSGGSITS
				PPLSPALPKYKLADYRYGREEMLALFLKDNKIPS
				DLLDKEFLPILQEEPLPPLALVPFTEEEQRNFSMS
				VNSAAVLRLTGRGGGGTVVGAPRGRSSSRGRGR
ļ		1		GRGECGFYQRSFDEVEGVFGRGGGREMHRSQS
		1		WEERGDRRFEKPGRKDVGRPNFEEGGPTSVGRK
		1		HEFIRSESENWRIFREEQNGEDEDGGWRLAGSRR
1	1	1	1	DGERWRPHSPDGPRSAGWREHMERRRFEFDFR
		9		DRDDERGYRRVRSGSGSIDDDRDSLPEWCLEDA
		1		EEEMGTFDSSGAFLSLKKVQKEPIPEEQEMDFRP
]		VDEGEECSDSEGSHNEEAKEPDKTNKKEGEKTD
1		1		RVGVEASEETPQTSSSSARPGTPSDHQSQEASQFE
				RKDEPKTEQTEKAEEETRMENSLPAKVPSRGDE
				MVADVQQPLSQIPSDTASPLLILPPPVPNPSPTLRP
		1		VETPVVGAPGMGSVSTEPDDEEGLKHLEQQAEK
<u> </u>	l	L	l	MVAYLQDSALDDERLASKLQEHRAKGVSIPLMH

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SEQ ID NO:	Method	Predicted beginning nucleotide location corresponding to first amino acid residue of peptide	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion
L		sequence		
		stquent		EAMQKWYYKDPQGEIQGPFNNQEMAEWFQAG YFTMSLLVKRACDESFQPLGDIMKMWGRVPFSP GPAPPPHMGELDQERLTRQQELTALYQMQHLQY QQFLIQQQYAQVLAQQQKAALSSQQQQQLALLL QQFQTLKMRISDQNIIPSVTRSVSVPDTGSIWELQ PTASQPTVWEGGSVWDLPLDTTTPGPALEQLQQ LEKAKAAKLEQERREAEMRAKREEEERKRQEEL RRRQKGILRRQQEEERKRREEEELARRKQEEALR RQREQEIALRRQREEEERQQQEEALRRLEERRRE EEERRKQEELLRKQEEEAAKWAREEEEAQRRLE ENRLRMEEEAARLRHEEEERKRKELEVQRQKEL MRQRQQQQEALRRLQQQQQQQQLAQMKLPSSS TWGQQSNTTACQSQATLSLAEIQKLEEERERQLR EEQRRQQRELMKALQQQQQQQQKLSGWGNV SKPSGTTKSLLEIQQEEARQMQKQQQQQQHQQ PNRARNNTHSNLHTSIGNSVWGSINTGPPNQWA SDLVSSIWSNADTKNSNMGFWDDAVKEVGPRN STNKNKNNASLSKSVGVSNRQNKKVEEEEKLLK LFQGVNKAQDGFTQWCEQMLHALNTANNLDVP TFVSFLKEVESPYEVHDYIRAYLGDTSEAKEFAK QFLERRAKQKANQQRQQQLPQQQQPPQQPP QQPQQQDSVWGMNHSTLHSVFQTNQSNNQQSN
				FEAVQSGKKKKKQKMVRADPSLLGFSVNASSER
2005 :		120	4050	LNMGEIETLDDY VERWINDERMA A ETOTI NECDEWI DAI SEGGSITS
3085		128		KSIVKIRKRMAAETQTLNFGPEWLRALSSGGSITS PPLSPALPKYKLADYRYGREEMLALFLKDNKIPS DLLDKEFLPILQEEPLPPLALVPFTEEEQRNFSMS VNSAAVLRLTGRGGGGTVVGAPRGRSSSRGRGR GRGECGFYQRSFDEVEGVFGRGGGREMHRSQS WEERGDRFEKPGRKDVGRPNFEEGGPTSVGRK HEFIRSESENWRIFREEQNGEDEDGGWRLAGSRR DGERWRPHSPDGPRSAGWREHMERRRFEFDFR DRDDERGYRRVRSGSGSIDDDRDSLPEWCLEDA EEEMGTFDSSGAFLSLKKVQKEPIPEEQEMDFRP VDEGEECSDSEGSHNEEAKEPDKTNKKEGEKTD RVGVEASEETPQTSSSSARPGTPSDHQSQEASQFE RKDEPKTEQTEKAEEETRMENSLPAKVPSRGDE MVADVQQPLSQIPSDTASPLLILPPPVPNPSPTLRP VETPVVGAPGMGSVSTEPDDEEGLKHLEQQAEK MVAYLQDSALDDERLASKLQEHRAKGVSIPLMH EAMQKWYYKDPQGEIQGPFNNQEMAEWFQAG YFTMSLLVKRACDESFQPLGDIMKMWGRVPFSP GPAPPPHMGELDQERLTRQQELTALYQMQHLQY QQFLIQQQYAQVLAQQQKAALSSQQQQLALLL QQFQTLKMRISDQNIIPSVTRSVSVPDTGSIWELQ PTASQPTVWEGGSVWDLPLDTTTPGPALEQLQQ LEKAKAAKLEQERREAEMRAKREEEELARRKQEEL RRQKGILRRQQEEERKRREEELARRKQEEALR RQREQEIALRRQREEEERQQCEALRRLEERRRE EEERRKQEELLRKQEEEAAKWAREEEEAQRRLE ENRLRMEEEAARLRHEEEERKRELEVQRQKEL MRQRQQQEALRRLQQQQQQQQLAQMKLPSSS TWGQQSNTTACQSQATLSLAEIQKLEEERRQLR EEQRRQQRELMKALQQQQQQQQQKLSGWGNV SKPSGTTKSLLEIQQEEARQMQKQQQQQQQHQQ

SEQ ID NO:	Method	Predicted beginning nucleotide location corresponding to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion
				PNRARNNTHSNLHTSIGNSVWGSINTGPPNQWA SDLVSSIWSNADTKNSNMGFWDDAVKEVGPRN STNKNKNNASLSKSVGVSNRQNKKVEEEEKLLK LFQGVNKAQDGFTQWCEQMLHALNTANNLDVP TFVSFLKEVESPYEVHDYIRAYLGDTSEAKEFAK QFLERRAKQKANQQRQQQLPQQQQPPQQPP QQPQQQDSVWGMNHSTLHSVFQTNQSNNQQSN FEAVQSGKKKKKQKMVRADPSLLGFSVNASSER LNMGEIETLDDY
3086	A	675	1334	LHPAATSTAWLHVPPGLSMALSWVLTVLSLLPL LEAQIPLCANLVPVPITNATLDRITGKWFYIASAF RNEEYNKSVQEIQATFFYFTPNKTEDTIFLREYQT RQDQCIYNTTYLNVQRENGTISRYVGGQEHFAH LLILRDTKTYMLAFDVNDEKNWGLSVYADKPET TKEQLGEFYEALDCLRIPKSDVVYTDWKKDKCE PLEKQHEKERKQEEGES
3087	A	1	1575	CTPVARSMATTATCTRFTDDYQLFEELGKGAFS VVRRCVKKTSTQEYAAKIINTKKLSARDHQKLE REARICRLLKHPNIVRLHDSISEEGFHYLVFDLVT GGELFEDIVAREYYSEADASHCIHQILESVNHIHQ HDIVHRDLKPENLLLASKCKGAAVKLADFGLAIE VQGEQQAWFGFAGTPGYLSPEVLRKDPYGKPVD IWACGVILYILLVGYPPFWDEDQHKLYQQIKAG AYDFPSPEWDTVTPEAKNLINQMLTINPAKRITA DQALKHPWVCQRSTVASMMHRQETVECLRKFN ARRKLKGAILTTMLVSRNFSAAKSLLNKKSDGG VKPQSNNKNSLVSPAQEPAPLQTAMEPQTTVVH NATDGIKGSTESCNTTTEDEDLKVRKQEIIKITEQ LIEAINNGDFEAYTKICDPGLTSFEPEALGNLVEG MDFHKFYFENLLSKNSKPIHTTILNPHVHVIGED AACIAYIRLTQYIDGQGRPRTSQSEETRVWHRRD GKWLNVHYHCSGAPAAPLQ
3088	A	12	1039	SSVAEFPERVQLSQPQNWNFSGAGGAWSLDFAE QLKWSAELARLGESIMDGKQGGMDGSKPAGPR DFPGIRLLSNPLMGDAVSDWSPMHEAAIHGHQL SLRNLISQGWAVNIITADHVSPLHEACLGGHLSC VKILLKHGAQVNGVTADWHTPLFNACVSGSWD CVNLLLQHGASVQPESDLASPIHEAARRGHVEC VNSLIAYGGNIDHKISHLGTPLYLACENQQRACV KKLLESGADVNQGKGQDSPLHAVARTASEELAC LLMDFGADTQAKNAEGKRPVELVPPESPLAQLF LEREGPPSLMQLCRLRIRKCFGIQQHHKITKLVLP EDLKQFLLHL
3089	A	73	432	DMAGLMTIVTSLLFLGVCAHHIIPTGSVVLPSPCC MFFVSKRIPENRVVSYQLSSRSTCLKAGVIFTTKK GQQFCGDPKQEWVQRYMKNLDAKQKKASPRA RAVAVKGPVQRYPGNQTTC
3090	A	4627	611	LMEAGGGGGALPAGVETMVLTLGESWPVLVGR RFLSLSAADGSDGSHDSWDVERVAEWPWLSGTI RAVSHTDVTKKDLKVCVEFDGESWRKRRWIEV YSLLRRAFLVEHNLVLAERKSPEISERIVQWPAIT YKPLLDKAGLGSITSVRFLGDQQRVFLSKDLLKP IQDVNSLRLSLTDNQIVSKEFQALIVKHLDESHLL KGDKNLVGSEVKIYSLDPSTQWFSATVVNGNPA SKTLQVNCEEIPALKIVDPSLIHVEVVHDNLVTC

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11 M WAINITACH

SEQ ID NO:	Method	Predicted beginning nucleotide location corresponding to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion
				GNSARIGA VKRKSSENNGTLVSKQAKSCSEASPS MCPVQSVPTTVFKEILLGCTAATPPSKDPRQQST PQAANSPPNLGAKIPQGCHKQSLPEEISSCLNTKS EALRTKPDVCKAGLLSKSSQIGTGDLKILTEPKGS CTQPKTNTDQENRLESVPQALTGLPKECLPTKAS SKAELEIANPPELQKHLEHAPSPSDVSNAPEVKA GVNSDSPNNCSGKKVEPSALACRSQNLKESSVK VDNESCCSRSNNKIQNAPSRKSVLTDPAKLKKLQ QSGEAFVQDDSCVNIVAQLPKCRECRLDSLRKD KEQQKDSPVFCRFFHFRRLQFNKHGVLRVEGFLT PNKYDNEAIGLWLPLTKNVVGIDLDTAKYILANI GDHFCQMVISEKEAMSTIEPHRQVAWKRAVKG VREMCDVCDTTIFNLHWVCPRCGFGVCVDCYR MKRKNCQQGAAYKTFSWLKCVKSQIHEPENLM PTQIIPGKALYDVGDIVHSVRAKWGIKANCPCSN RQFKLFSKPASKEDLKQTSLAGEKPTLGAVLQQ NPSVLEPAAVGGEAASKPAGSMKPACPASTSPLN WLADLTSGNVNKENKEKQPTMPILKNEIKCLPPL PPLSKSSTVLHTFNSTILTPVSNNNSGFLRNLLNSS TGKTENGLKNTPKILDDIFASLVQNKTTSDLSKR PQGLTIKPSILGFDTPHYWLCDNRLLCLQDPNNK SNWNVFRECWKQGQPVMVSGVHHKLNSELWK PESFRKEFGEQEVDLVNCRTNEIITGATVGDFWD GFEDVPNRLKNEKEPMVLKLKDWPPGEDFRDM MPSRFDDLMANIPLPEYTRRDGKLNLASRLPNYF VRPDLGPKMYNAYGLITPEDRKYGTTNLHLDVS DAANVMVYVGIPKGQCEQEEEVLKTIQDGDSDE LTIKRFIEGKEKPGALWHIYAAKDTEKIREFLKK VSEEQGQENPADHDPIHDQSWYLDRSLRKRLHQ EYGVQGWAIVQFLGDVVFIPAGAPHQVHNLYSC IKVAEDFVSPEHVKHCFWLTQEFRYLSQTHTNHE DKLQVKNVIYHAVKDAVAMLKASESSFGKP
3091	A	97	1838	KRGARRGGWKRKMPSTDLLMLKAFEPYLEILEV YSTKAKNYVNGHCTKYEPWQLIAWSVVWTLLI VWGYEFVFQPESLWSRFKKKCFKLTRKMPIIGRK IQDKLNKTKDDISKNMSFLKVDKEYVKALPSQG LSSSAVLEKLKEYSSMDAFWQEGRASGTVYSGE EKLTELLVKAYGDFAWSNPLHPDIFPGLRKIEAEI VRIACSLFNGGPDSCGCVTSGGTESILMACKAYR DLAFEKGIKTPEIVAPQSAHAAFNKAASYFGMKI VRVPLTKMMEVDVRAMRRAISRNTAMLVCSTP QFPHGVIDPVPEVAKLAVKYKIPLHVDACLGGFL IVFMEKAGYPLEHPFDFRVKGVTSISADTHKYGY APKGSSLVLYSDKKYRNYQFFVDTDWQGGIYAS PTIAGSRPGGISAACWAALMHFGENGYVEATKQI IKTARFLKSELENIKGIFVFGNPQLSVIALGSRDFD IYRLSNLMTAKGWNLNQLQFPPSIHFCITLLHAR KRVAIQFLKDIRESVTQIMKNPKAKTTGMGAIYG MAQTTVDRNMGAELSSVFLDSLYSTDTVTQGSQ MNGSPKPH
3092	A	79	2652	LCSQNSPEDWVNFSSEKQKRYPWYWTGRKLRSE RAMKIQKKLTGCSRLMLLCLSLELLLEAGAGNIH YSVPEETDKGSFVGNIAKDLGLQPQELADGGVRI VSRGRMPLFALNPRSGSLITARRIDREELCAQSM PCLVSFNILVEDKMKLFPVEVEIIDINDNTPQFQL

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	·			EELEFKMNEITTPGTRVSLPFGQDLDVGMNSLQS YQLSSNPHFSLDVQQGADGPQHPEMVLQSPLDR EEEAVHHLILTASDGGEPVRSGTLRIYIQVVDAN DNPPAFTQAQYHINVPENVPLGTQLLMVNATDP DEGANGEVTYSFHNVDHRVAQIFRLDSYTGEISN KEPLDFEEYKMYSMEVQAQDGAGLMAKVKVLI KVLDVNDNAPEVTITSVTTAVPENFPPGTIIALISV HDQDSGDNGYTTCFIPGNLPFKLEKLVDNYYRL VTERTLDRELISGYNITITAIDQGTPALSTETHISL LVTDINDNSPVFHQDSYSAYIPENNPRGASIFSVR AHDLDSNENAQITYSLIEDTIQGAPLSAYLSINSD TGVLYALRSFDYEQFRDMQLKVMARDSGDPPLS SNVSLSLFLLDQNDNAPEILYPALPTDGSTGVEL APRSAEPGYLVTKVVAVDRDSGQNAWLSYRLL KASEPGLFSVGLHTGEVRTARALLDRDALKQSL VVAVQDHGQPPLSATVTLTVAVADRIPDILADLG SLEPSAKPNDSDLTLYLVVAEAAVSCVFLAFVIV LLAHRLRRWHKSRLLQASGGGLASTPGSHFVGV DGVRAFLQTYSHEVSLTADSRKSHLIFPQPNYAD TLISQESCEKKGFLSAPQSLLEDKKEPFSQVNFCD ECISYLEKNNS
3093	A	1	3868	PPDNQKLGLLEALLKIGDWQHAQNIMDQMPPYY AASHKLIALAICKLIHITIEPLYRSVTSWAVDHAG FLESDPCDSTVGHLLSRVGVPKGAKGSPVNALQ NKRAPKQAESFEDLRDVFNMFCYLGPHLSHDPI LFAKVVRIGKSFMKEFQSDGSKQEDKEKTEVILS CLLSITDQVLLPSLSLMDCNACMSEELWGMFKT FPYQHRYRLYGQWKNETYNSHPLLVKVKAQTID RAKYIMKRLTKENVKPSGRQIGKLSHSNPTILFD YVCFEILSQIQKYDNLITPVVDSLKYLTSLNYDVL ACILSNCIIEALANPEKERMKHDDTTISSWLQSLA SFCGAVFRKYPIDLAGLLQYVANQLKAGKSFDL LILKEVVQKMAGIEITEEMTMEQLEAMTGGEQL KAEGGYFGQIRNTKKSSQRLKDALLDHDLALPL CLLMAQQRNGVIFQEGGEKHLKLVGKLYDQCH DTLVQFGGFLASNLSTEDYIKRVPSIDVLCNEFHT PHDAAFFLSRPMYAHHISSKYDELKKSEKGSKQ QHKVHKYITSCEMVMAPVHEAVVSLHVSKVWD DISPQFYATFWSLTMYDLAVPHTSYEREVNKLK VQMKAIDDNQEMPPNKKKKEKERCTALQDKLL EEEKKQMEHVQRVLQRLKLEKDNWLLAKSTKN ETITKFLQLCIFPRCIFSAIDAVYCARFVELVHQQ KTPNFSTLLCYDRVFSDIIYTVASCTENEASRYGR FLCCMLETVTRWHSDRATYEKECGNYPGFLTIL RATGFDGGNKADQLDYENFRHVVHKWHYKLT KASVHCLETGEYTHIRNILIVLTKILPWYPKVLNL GQALERRVHKICQEEKEKRPDLYALAMGYSGQL KSRKSYMIPENEFHHKDPPPRNAVASVQNGPGG GPSSSSIGSASKSDESSTEETDKSRERSQCGVKAV NKASSTTPKGNSSNGNSGSNSNKAVKENDKEKG KEKEKEKEKEKTPATTPEARVLGKDGKEKPKEER PNKDEKARETKERTPKSDKEKEKFKKEEKAKDE KFKTTVPNAESKSTQEREREKEPSRERDIAKEMK SKENVKGGEKTPVSGSLKSPVPRSDIPEPEREQKR RKIDTHPSPSHSSTVKDSLIELKESSAKLYNHTPP

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		-		PLSKSKEREMDKKDLDKSRERSREREKKDEKDR KERKRDHSNNDREVPPDLTKRRKEENGTMGVSK HKSESPCESPYPNEKDKEKNKSKSSGKEKGSDSF KSEKMDKISSGGKKESRHDKEKIEKKEKRDSSGG KEEKKHHKSSDKHR
3094		2	891	AMLGTREPSRRGAGAVQAEVSERLAMAGPQQQ PPYLHLAELTASQFLEIWKHFDADGNGYIEGKEL ENFFQELEKARKGSGMMSKSDNFGEKMKEFMQ KYDKNSDGKIEMAELAQILPTEENFLLCFRQHVG SSAEFMEAWRKYDTDRSGYIEANELKGFLSDLL KKANRPYDEPKLQEYTQTILRMFDLNGDGKLGL SEMSRLLPVQENFLLKFQGMKLTSEEFNAIFTFY DKDRSGYIDEHELDALLKDLYEKNKKEMNIQQL TNYRKSVMSLAEAGKLYRKDLEIVLCSEPPM
3095	A	1685	700	RRPTGRPGALGAPAAGRVGMPLHVKWPFPAVPP LTWTLASSVVMGLVGTYSCFWTKYMNHLTVHN REVLYELIEKRGPATPLITVSNHQSCMDDPHLWG ILKLRHIWNLKLMRWTPAAADICFTKELHSHFFS LGKCVPVCRGAEFFQAENEGKGVLDTGRHMPG AGKRREKGDGVYQKGMDFILEKLNHGDWVHIF PEGKVNMSSEFLRFKWGIGRLIAECHLNPIILPLW HVGMNDVLPNSPPYFPRFGQKITVLIGKPFSALP VLERLRAENKSAVEMRKALTDFIQEEFQHLKTQ AEQLHNHLQAWEIGLACCLLDSWPAQSWG
3096	A	6642	4022	FVPGLREPQWEPAQPSATMSAPSEEEEYARLVM EAQPEWLRAEVKRLSHELAETTREKIQAAEYGL AVLEEKHQLKLQFEELEVDYEAIRSEMEQLKEAF GQAHTNHKKVAADGESREESLIQESASKEQYYV RKVLELQTELKQLRNVLTNTQSENERLASVAQE LKEINQNVEIQRGRLRDDIKEYKFREARLLQDYS ELEEENISLQKQVSVLRQNQVEFEGLKHEIKRLE EETEYLNSQLEDAIRLKEISERQLEEALETLKTER EQKNSLRKELSHYMSINDSFYTSHLHVSLDGLKF SDDAAEPNNDAEALVNGFEHGGLAKLPLDNKTS TPKKEGLAPPSPSLVSDLLSELNISEIQKLKQQLM QMEREKAGLLATLQDTQKQLEHTRGSLSEQQEK VTRLTENLSALRRLQASKERQTALDNEKDRDSH EDGDYYEVDINGPEILACKYHVAVAEAGELREQ LKALRSTHEAREAQHAEEKGRYEAEGQALTEKV SLLEKASRQDRELLARLEKELKKVSDVAGETQG SLSVAQDELVTFSEELANLYHHVCMCNNETPNR VMLDYYREGQGGAGRTSPGGRTSPEARGRRSPI LLPKGLLAPEAGRADGGTGDSSPSPGSSLPSPLSD PRREPMNIYNLIAIIRDQIKHLQAAVDRTTELSRQ RIASQELGPAVDKDKEALMEEILKLKSLLSTKRE QITTLRTVLKANKQTAEVALANLKSKYENEKAM VTETMMKLRNELKALKEDAATFSSLRAMFATRC DEYITQLDEMQRQLAAAEDEKKTLNSLLRMAIQ QKLALTQRLELLELDHEQTRRGRAKAAPKTKPA TPSVSHTCACASDRAEGTGLANQVFCSEKHSIYC D
3097	A	1	879	MVKVVPATRGNLPRSQLTGTHQHCQPREPKITA SERLRRPRATARLRAHAAPPEPPLAVFAPPSDR KELLALPVACDPVIASVMSWVQAASLIQGPGDK GDVFDEEADESLLAQREWQSNMQRRVKEGYRD

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NO:		beginning nucleotide	nucleotide location	I=Isoleucine, K=Lysine, L=Leucine, M=Methionine,
		location	corresponding	N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine,
		corresponding to first amino	to last amino acid residue of	T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion,
		acid residue of	peptide	-possible nucleotide insertion
		peptide sequence	sequence	
				GIDAGKAVTLQQGFNQGYKKGAEVILNYGRLRG TLSALLSWCHLHNNNSTLINKINNLLDAVGQCEE
				YVLKHLKSITPPSHVVDLLDSIEDMDLCHVVPAE
•		1		KKIDEAKDERLCENNAEFNKNCSKSHSGIDCSYV
		1		ECCRTQEHAHSGKPKPHMDFGTDSQF
3098	Α	2	505	GAATLLRSASSAARKAAEAEQVWLHLHRYLSA
	1	İ	ļ	DRRVLGLREWGRPASERECSLCQRLKRELNMGD
				VEKGKKIFIMKCSQCHTVEKGGKHKTGPNLHGL
				FGRKTGQAPGYSYTAANKNKGIIWGEDTLMEYL ENPKKYIPGTKMIFVGIKKKEERADLIAYLKKAT
	İ			NE
3099	A	144	1386	WAVGQARSFPSHPRMSSWIWSRRWSPSVALRVT
				CTSTSSQRWTVLALSKPGSQQQVSMHTPAPGPPT
		1		AGHTEPPSEPPRRARVAKYRAKFDPRVTAKYDIK
	i		i .	ALIGRGSFSRVVRVEHRATRQPYAIKMIETKYRE GREVCESELRVLRRVRHANIIQLVEVFETQERVY
	1			MVMELATGGELFDRIIAKGSFTERDATRVLQMV
				LDGVRYLHALGITHRDLKPENLLYYHPGTDSKIII
	1	}	}	TDFGLASARKKGDDCLMKTTCGTPEYIAPEVLV
				RKPYTNSVDMWALGVIAYILLSGTMPFEDDNRT
	1		•	RLYRQILRGKYSYSGEPWPSVSNLAKDFIDRLLT
]			VDPGARMTALQALRHPWVVSMAASSSMKNLHR
	1			SISQNLLKRASSRCQSTKSAQSTRSSRSTRSNKSR RVRERELREL
3100	A	3	1500	ARWNGRWVQVPAWPGPGCGTNASGERQRQLPR
3100	A		1500	AWRPVGRTLGSEPIALAWSPPLYLFPIPLPSWAVS
				QPTPTLGTMFADLDYDIEEDKLGIPTVPGKVTLQ
				KDAQNLIGISIGGGAQYCPCLYIVQVFDNTPAAL
			ł	DGTVAAGDEITGVNGRSIKGKTKVEVAKMIQEV
)	KGEVTIHYNKLQADPKQGMSLDIVLKKVKHRLV ENMSSGTADALGLSRAILCNDGLVKRLEELERTA
				ELYKGMTEHTKNLLRAFYELSQTHRGNGIPQSC
				AFGDVFSVIGVREPQPAASEAFVKFADAHRSIEK
				FGIRLLKTIKPMLTDLNTYLNKAIPDTRLTIKKYL
				DVKFEYLSYCLKVKEMDDEEYSCIALGEPLYRV
				STGNYEYRLILRCRQEARARFSQMRKDVLEKME
				LLDQKHVQDIVFQLQRLVSTMSKYYNDCYAVLR
				DADVFPIEVDLAHTTLAYGLNQEEFTDGEEEEEE EDTAAGEPSRDTRGAAGPLDKGGSWCDS
3101	A	1173	197	OGMDSKQQCVKLNDGHFMPVLGFGTYAPPEVP
				RSKALEVTKLAIEAGFRHIDSAHLYNNEEQVGLA
				IRSKIADGSVKREDIFYTSKLWSTFHRPELVRPAL
			[ENSLKKAQLDYVDLYLIHSPMSLKPGEELSPTDE
				NGKVIFDIVDLCTTWEAMEKCKDAGLAKSIGVS
				NFNRRQLEMILNKPGLKYKPVCNQVECHPYFNR SKLLDFCKSKDIVLVAYSALGSQRDKRWVDPNS
	}	1		PVLLEDPVLCALAKKHKRTPALIALRYQLQRGV
				VVLAKSYNEQRIRQNVQVFEFQLTAEDMKAIDG
				LDRNLHYFNSDSFASHPNYPYSDEY
		1		EOPRPPPCGRRPLPLGSAPCRVRLGRAPRQAPAM
3102	A	144	1098	
3102	A	144	1098	SMLPSFGFTQEQVACVCEVLQQGGNLERLGRFL
3102	A	144	1098	SMLPSFGFTQEQVACVCEVLQQGGNLERLGRFL WSLPACDHLHKNESVLKAKAVVAFHRGNFREL
3102	A	144	1098	SMLPSFGFTQEQVACVCEVLQQGGNLERLGRFL WSLPACDHLHKNESVLKAKAVVAFHRGNFREL YKILESHQFSPHNHPKLQQLWLKAHYVEAEKLR
3102	A	144	1098	SMLPSFGFTQEQVACVCEVLQQGGNLERLGRFL WSLPACDHLHKNESVLKAKAVVAFHRGNFREL

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		ē		TQVSNWFKNRRQRDRAAEAKERENTENNNSSSN KQNQLSPLEGGKPLMSSSEEEFSPPQSPDQNSVLL LQGNMGHARSSNYSLPGLTASQPSHGLQTHQHQ LQDSLLGPLTSSLVDLGS
3103	A	111	1582	LVYSWGCHIMADNDTDRNQTEKLLKRVRELEQ EVQRLKKEQAKNKEDSNIRENSSGAGKTKRAFD FSAHGRRHVALRIAYMGWGYQGFASQENTNNTI EEKLFEALTKTRLVESRQTSNYHRCGRTDKGVS AFGQVISLDLRSQFPRGRDSEDFNVKEEANAAAE EIRYTHILNRVLPPDIRILAWAPVEPSFSARFSCLE RTYRYFFPRADLDIVTMDYAAQKYVGTHDFRNL CKMDVANGVINFQRTILSAQVQLVGQSPGEGRW QEPFQLCQFEVTGQAFLYHQVRCMMAILFLIGQ GMEKPEIIDELLNIEKNPQKPQYSMAVEFPLVLY DCKFENVKWIYDQEAQEFNITHLQQLWANHAV KTHMLYSMLQGLDTVPVPCGIGPKMDGMTEWG NVKPSVIKQTSAFVEGVKMRTYKPLMDRPKCQG LESRIQHFVRRGRIEHPHLFHEEETKAKRDCNDT LEEDNTNLETPTKRVCVDTEIKSII
3104	A	227	1519	VTLIKMNAMLETPELPAVFDGVKLAAVAAVLYV IVRCLNLKSPTAPPDLYFQDSGLSRFLLKSCPLLT KEYIPPLIWGKSGHIQTALYGKMGRVRSPHPYGH RKFITMSDGATSTFDLFEPLAEHCVGDDITMVICP GIANHSEKQYIRTFVDYAQKNGYRCAVLNHLGA LPNIELTSPRMFTYGCTWEFGAMVNYIKKTYPLT QLVVVGFSLGGNIVCKYLGETQANQEKVLCCVS VCQGYSALRAQETFMQWDQCRRFYNFLMADN MKKIILSHRQALFGDHVKKPQSLEDTDLSRLYTA TSLMQIDDNVMRKFHGYNSLKEYYEEESCMRYL HRIYVPLMLVNAADDPLVHESLLTIPKSLSEKRE NVMFVLPLHGGHLGFFEGSVLFPEPLTWMDKLV VEYANAICQWERNKLQCSDTEQVEADLE
3105	A		1251	MGLLMILASAVLGSFLTLLAQFFLLYRRQPEPP ADEAARAGEGFRYIKPVPGLLLREYLYGGGRDE EPSGAAPEGGATPTAAPETPAPPTRETCYFLNATI LFLFRELRDTALTRRWVTKKIKVEFEELLQTKTA GRLLEGLSLRDVFLGETVPFIKTIRLVRPVVPSAT GEPDGPEGEALPAACPEELAFEAEVEYNGGFHLA IDVDLVFGKSAYLFVKLSRVVGRLRLVFTRVPFT HWFFSFVEDPLIDFEVRSQFEGRPMPQLTSIIVNQ LKKIIKRKHTLPNYKIRFKPFFPYQTLQGFEEDEE HIHIQQWALTEGRLKVTLLECSRLLIFGSYDREA NVHCTLELSSSVWEEKQRSSIKTGTISLTAVFMG WHRVSEAFPGLWYKLLVDLPFWGLEDGGPLLT VPLRQCPG
3106	A	972	468	MAAAGAGRLRRVASALLLRSPRLPARELSAPAR LYHKKVVDHYENPRNVGSLDKTSKNVGTGLVG APACGDVMKLQIQVDEKGKIVDARFKTFGCGSA IASSSLATEWVKGKTVEEALTIKNTDIAKELCLPP VKLHCSMLAEDAIKAALADYKLKQEPKKGEAE KK
3107	A	106	1221	TCQDVRSVFSLVRANIFGEESTAGAGWHREEDM RKELQLSLSVTLLLVCGFLYQFTLKSSCLFCLPSF KSHQGLEALLSHRRGIVFLETSERMEPPHLVSCS VESAAKIYPEWPVVFFMKGLTDSTPMPSNSTYPA

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				FSFLSAIDNVFLFPLDMKRLLEDTPLFSWYNQINA SAERNWLHISSDASRLAIIWKYGGIYMDTDVISIR PIPEENFLAAQASRYSSNGIFGFLPHHPFLWECME NFVEHYNSAIWGNQGPELMTRMLRVWCKLEDF QEVSDLRCLNISFLHPQRFYPISYREWRRYYEVW DTEPSFNVSYALHLWNHMNQEGRAVIRGSNTLV ENLYRKHCPRTYRDLIKGPEGSVTGELGPGNK
3108	A	1612	839	EVALFCFEMAAGMYLEHYLDSIENLPFELQRNFQ LMRDLDQRTEDLKAEIDKLATEYMSSARSLSSEE KLALLKQIQEAYGKCKEFGDDKVQLAMQTYEM VDKHIRRLDTDLARFEADLKEKQIESSDYDSSSS KGKKKGRTQKEKKAARARSKGKNSDEEAPKTA QKKLKLVRTSPEYGMPSVTFGSVHPSDVLDMPV DPNEPTYCLCHQVSYGEMIGCDNPDCSIEWFHFA CVGLTTKPRGKWFCPRCSQERKKK
	A		2613	MVAVRAAGPREGASQDEAGTVWAPMTGCPCQC RPGPSWLLVDTLEPETAYPVQRPGPEQAGNQRL QMKRAQFGPHDWLSLPVPPGPSWLLVDTLEPET AYQFSVLAQNKLGTSAFSEVVTVNTLAFPITTPEP LVLVTPPRCLIANRTQQGVLLSWLPPANHSFPIDR YIMEFRVAERWELLDDGIPGTEGEFFAKDLSQDT WYEFRVLAVMQDLISEPSNIAGVSSTDIFPQPDLT EDGLARPVLAGIVATICFLAAAILFSTLAACFVNK QRKRKLKRKKDPPLSITHCRKSLESPLSSGKVSPE SIRTLRAPSESSDDQGQPAAKRMLSPTREKELSL YKKTKRAISSKKYSVAKAEAEAEATTPIELISRGP DGRFVMDPAEMEPSLKSRRIEGFPFAEETDMYPE FRQSDEENEDPLVPTSVAALKSQLTPLSSSQESYL PPPAYSPRFQPRGLEGPGGLEGRLQATGQARPPA PRPFHHGQYYGYLSSSSPGEVEPPPFYVPEVGSPL SSVMSSPPLPTEGPFGHPTIPEENGENASNSTLPLT QTPTGGRSPEPWGRPEFPFGGLETPAMMFPHQLP PCDVPESLQPKAGLPRGLPPTSLQVPAAYPGILSL EAPKGWAGKSPGRGPVPAPPAAKWQDRPMQPL VSQGQLRHTSQGMGIPVLPYPEPAEPGAHGGPST FGLDTRWYEPQPRPRSPRQARRAEPSLHQVVLQ PSRLSPLTQSPLSSRTGSPELAARARPRPGLLQQA EMSEITLQPPAAVSFSRKSTPSTGSPSQSSRSGSPS YRPAMGFTTLATGYPSPPPGPAPAGPGDSLDVFG QTPSPRRTGEELLRPETPPPTLPTLGKLRRDRPAP ATSPPERALSKL
3110	A	88	924	ILGSRTMSLTNTKTGFSVKDILDLPDTNDEEGSV AEGPEEENEGPEPAKRAGPLGQGALDAVQSLPL KNPFYDSSDNPYTRWLASTEGLQYSLHGLAAGA PPQDSSSKSPEPSADESPDNDKETPGGGGDAGKK RKRRVLFSKAQTYELERRFRQQRYLSAPEREHLA SLIRLTPTQVKIWFQNHRYKMKRARAEKGMEVT PLPSPRRVAVPVLVRDGKPCHALKAQDLAAATF QAGIPFSAYSAQSLQHMQYNAQYSSASTPQYPT AHPLVQAQQWTW
3111	A	595 3641	291 . 1555	PSVASLARRFSGRALWPPSHSVPGNRALCPRLLH GTTLPGGNQRELARQKNMKKQSDSVKGKRRDD GLSAAARKQRDSTPRDSEIMQQKQKKANEKKEE PK APMLQIHHFSFKLIFQNIHKSKFISQRLSQNADST

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·				RHTNLSNTHYSDLIVWNCCLFFRNWCNEFFLKS CHFAQEREGSGDLCNSRAEKTKSAACVIFRRFPV APLIPYPLITKEDINAIEMEEDKRDLISREISKFRDT HKKLEEEKGKKEKERQEIEKERRERERERERERE RREREREREREREREKEKERERERER
3113	A	1 · · · · · · · · · · · · · · · · · · ·	669	LLIYETEAKKIGLVK VCAGIRDPCSTPLAKPAAGGAENLSFGKQPGLET NILKMTTPNKTPPGADPKQLERTGTVREIGSQAV WSLSSCKPGFGVDQLRDDNLETYWQSDGSQPHL VNIQFRRKTTVKTLCIYADYKSDESYTPSKISVRV GNNFHNLQEIRQLELVEPSGWIHVPLTDNHKKPT RTFMIQIAVLANHQNGRDTHMRQIKIYTPVEESSI GKFPRCTTIDFMMYRSIR
3114	A		1613	MTSKEESRRQQPTAGPAGQGKLPSPSEPQLPTPP TRSLHHFRRPLSPSREAQAHIAPSSELHLPQSQSA GPPPLGAGTEVELVVPGRDEGSRGALPGSSGVKF VWRKIVRFPVSDQVRTLSISRLMRRLLEMMQTL VQFIIGWRSLLGRTLGTIMNTMYVMMAQILRSH LIKATVIPNRVKMLPYFGIIRNRMMSTHKSKKKI REYYRLLNVEEGCSADEVRESFHKLAKQYHPDS GSNTADSATFIRIEKAYRKVLSHVIEQTNASQSK GEEEDVEKFKYKTPQHRHYLSFEGIGFGTPTQR EKHYRQFRADRAAEQVMEYQKQKLQSQYFPDS VIVKNIRQSKQQKITQAIERLVEDLIQESMAKGDF DNLSGKGKPLKKFSDCSYIDPMTHNLNRILIDNG YQPEWILKQKEISDTIEQLREAILVSRKKLGNPMT PTEKKQWNHVCEQFQENIRKLNKRINDFNLIVPI LTRQKVHFDAQKEIVRAQKIYETLIKTKEVTDRN PNNLDQGEGEKTPEIKKGFLNLMDLVEIY
3115	A		2036	FRHRCGCLSYCRSRRGIRRVEPLRRARARVGPRF RPLCRMEIIRSNFKSNLHKVYQAIEEADFFAIDGE FSGISDGPSVSALTNGFDTPEERYQKLKKHSMDF LLFQFGLCTFKYDYTDSKYITKSFNFYVFPKPFNR SSPDVKFVCQSSSIDFLASQGFDFNKGFRKGIPYL NQEEERQLREQYDEKRSQANGAGALSYVSPNTS KCPVTIPEDQKKFIDQVVEKIEDLLQSEENKNLDL EPCTGFQRKLIYQTLSWKYPKGIHVETLETEKKE RYIVISKVDEEERKRREQQKHAKEQEELNDAVG FSRVIHAIANSGKLVIGHNMLLDVMHTVHQFYC PLPADLSEFKEMTTCVFPRLLDTKLMASTQPFKD IINNTSLAELEKRLKETPFNPPKVESAEGFPSYDT

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				ASEQLHEAGYDAYITGLCFISMANYLGSFLSPPKI HVSARSKLIEPFFNKLFLMRVMDIPYLNLEGPDL QPKRDHVLHVTFPKEWKTSDLYQLFSAFGNIQIS WIDDTSAFVSLSQPEQVKIAVNTSKYAESYRIQT YAEYMGRKQEEKQIKRKWTEDSWKEADSKRLN PQCIPYTLQNHYYRNNSFTAPSTVGKRNLSPSQE EAGLEDGVSGEISDTELEQTDSCAEPLSEGRKKA KKLKRMKKELSPAGSISKNSPATLFEVPDTW
3116	A	3	1443	TREAPMALAVAPWGRQWEEARALGRAVRMLQ RLEEQCVDPRLSVSPPSLRDLLPRTAQLLREVAH SRRAAGGGPGPGPGGSGDFLLIYLANLEAKSRQ VAALLPPRGRRSANDELFRAGSRLRRQLAKLAII FSHMHAELHALFPGGKYCGHMYQLTKAPAHTF WRESCGARCVLPWAEFESLLGTCHPVEPGCTAL ALRTTIDLTCSGHVSIFEFDVFTRLFQPWPTLLKN WQLLAVNHPGYMAFLTYDEVQERLQACRDKPG SYIFRPSCTRLGQWAIGYVSSDGSILQTIPANKPLS QVLLEGQKDGFYLYPDGKTHNPDLTELGQAEPQ QRIHVSEEQLQLYWAMDSTFELCKICAESNKDV KIEPCGHLLCSCCLAAWQHSDSQTCPFCRCEIKG WEAVSIYQFHGQATAEDSGNSSDQEGRELELGQ VPLSAPPLPPRPDLPPRKPRNAQPKVRLLKGNSPP AALGPQDPAPA
3117	A	296	3547	ERHSSPLLQHILTHALMRNKKHSNNWLAQHWF QSSIILCFSPVGRTLRVRARKFPAIVNCTAIDWFH AWPQEALVSVSRRFIEETKGIEPVHKDSISLFMAH VHTTVNEMSTRYYQNERRHNYTTPKSFLEQISLF KNLLKKKQNEVSEKKERLVNGIQKLKTTASQVG DLKARLASQEAELQLRNHDAEALITKIGLQTEKV SREKTIADAEERKVTAIQTEVFQKQRECEADLLK AEPALVAATAALNTLNRVNLSELKAFPNPPIAVT NVTAAVMVLLAPRGRVPKDRSWKAAKVFMGK VDDFLQALINYDKEHIPENCLKVVNEHYLKDPEF NPNLIRTKSFAAAGLCAWVINIIKFYEVYCDVEP KRQALAQANLELAAATEKLEAIRKKLVVSANYD IEKSEKIRWGQSIKSFEAQEKTLCGDVLLTAAFVS YVGPFTRQYRQELVHCKWVPFLQQKVSIPLTEG LDLISMLTDDATIAAWNNEGLPSDRMSTENAAIL THCERWPLVIDPQQQGIKWIKNKYGMDLKVTHL GQKGFLNAIETALAFGDVILIENLEETIDPVLDPL LGRNTIKKGKYIRIGDKECEFNKNFRLILHTKLAN PHYKPELQAQTTLLNFTVTEDGLEAQLLAEVVSI ERPDLEKLKLVLTKHQNDFKIELKYLEDDLLLRL SAAEGSFLDDTKLVERLEATKTTVAEIEHKVIEA KENERKINEARECYRPVAARASLLYFVINDLQKI NPLYQFSLKAFNVLFHRAIEQADKVEDMQGRISI LMESITHAVFLYTSQALFEKDKLTFLSQMAFQIL LRKKEIDPLELDFLLRFTVEHTHLSPVDFLTSQSW SAIKAIAVMEEFRGIDRDVEGSAKQWRKWVESE CPEKEKLPQEWKKKSLIQKLILLRAMRPDRMTY ALRNFVEEKLGAKYVERTRLDLVKAFEESSPATP IFFILSPGVDALKDLEILGKRLGFTIDSGKFHNVSL GQGQETVAEVALEKASKGGHWVILQNVHLVAK WLGTLEKLLERFSQGSHRDYRVFMSAESAPTPD EHIIPQGLLENSIKITNEPPTGMLANLHAALYNFD

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3118	A	1	226	Q PYSLSTSCLGSPTSPRLEMDPNCSCATGGSCTCTG SCKCKECKCNSCKKSECGAISRNLGLSQVRGRKP ELGMEE
3119	A	1254	4133	PLATLTMEEQGHSEMEIIPSESHPHIQLLKSNREL LVTHIRNTQCLVDNLLKNDYFSAEDAEIVCACPT QPDKVRKILDLVQSKGEEVSEFFLYLLQQLADAY VDLRPWLLEIGFSPSLLTQSKVVVNTDPVSRYTQ QLRHHLGRDSKFVLCYAQKEELLLEEIYMDTIME LVGFSNESLGSLNSLACLLDHTTGILNEQGETIFIL GDAGVGKSMLLQRLQSLWATGRLDAGVKFFFH FRCRMFSCFKESDRLCLQDLLFKHYCYPERDPEE VFAFLLRFPHVALFTFDGLDELHSDLDLSRVPDS SCPWEPAHPLVLLANLLSGKLLKGASKLLTART GIEVPRQFLRKKVLLRGFSPSHLRAYARRMFPER ALQDRLLSQLEANPNLCSLCSVPLFCWIIFRCFQH FRAAFEGSPQLPDCTMTLTDVFLLVTEVHLNRM QPSSLVQRNTRSPVETLHAGRDTLCSLGQVAHR GMEKSLFVFTQEEVQASGLQERDMQLGFLRALP ELGPGGDQQSYEFFHLTLQAFFTAFFLVLDDRVG TQELLRFFQEWMPPAGAATTSCYPPFLPFQCLQG SGPAREDLFKNKDHFQFTNLFLCGLLSKAKQKLL RHLVPAAALRRKRKALWAHLFSSLRGYLNSLPR VQVESFNQVQAMPTFIWMLRCIYETQSQKVGQL AARGICANYLKLTYCNACSADCSALSFVLHHFP KRLALDLDNNNLNDYGVRELQPCFSRLTVLRLS VNQITDGGVKVLSEELTKYKIVTYLGLYNNQITD VGARYVTKILDECKGLTHLKLGKNKITSEGGKY LALAVKNSKSISEVGMWGNQVGDEGAKAFAEA LRNHPSLTTLSLASNGISTEGGKSLARALQQNTSL EILWLTQNELNDEVAESLAEMLKVNQTLKHLWL IQNQITAKGTAQLADALQSNTGITEICLNGNLIKP
3120	A	43	1004	EEAKVYEDEKRIICF QLWGFAAGSDSRPAMGCDGGTIPKRHELVKGPK KVEKVDKDAELVAQWNYCTLSQEILRRPIVACE LGRLYNKDAVIEFLLDKSAEKALGKAASHIKSIK NVTELKLSDNPAWEGDKGNTKGDKHDDLQRAR FICPVVGLEMNGRHRFCFLRCCGCVFSERALKEI KAEVCHTCGAAFQEDDVIVLNGTKEDVDVLKTR MEERRLRAKLEKKTKKPKAAESVSKPDVSEEAP GPSKVKTGKPEEASLDSREKKTNLAPKSTAMNE SSSGKAGKPPCGATKRSIADSEESEAYKSLFTTHS SAKRSKEESAHWVTHTSYCF
3121	A	3	1490	HASGPTRPVSWSFHKLKTMKHLLLLLLCVFLVK SQGVNDNEEGFFSARGHRPLDKKREEAPSLRPAP PPISGGGYRARPAKAAATQKKVERKAPDAGGCL HADPDLGVLCPTGCQLQEALLQQERPIRNSVDEL NNVEAVSQTSSSSFQYMYLLKDLWQKRQKQV KDNENVVNEYSSELEKHQLYIDETVNSNIPTNLR VLRSILENLRSKIQKLESDVSAQMEYCRTPCTVS CNIPVVSGKECEEIIRKGGETSEMYLIQPDSSVKP YRVYCDMNTENGGWTVIQNRQDGSVDFGRKW DPYKQGFGNVATNTDGKNYCGLPGEYWLGNDK ISQLTRMGPTELLIEMEDWKGDKVKAHYGGFTV QNEANKYQISVNKYRGTAGNALMDGASQLMGE

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				NRTMTIHNGMFFSTYDRDNDGWLTSDPRKQCSK EDGGGWWYNRCHAANPNGRYYWGGQYTWDM AKHGTDDGVVWMNWKGSWYSMKKMSMKIRP FFPQQ
3122	A	3	1490	HASGPTRPVSWSFHKLKTMKHLLLLLLCVFLVK SQGVNDNEEGFFSARGHRPLDKKREEAPSLRPAP PPISGGGYRARPAKAAATQKKVERKAPDAGGCL HADPDLGVLCPTGCQLQEALLQQERPIRNSVDEL NNNVEAVSQTSSSSFQYMYLLKDLWQKRQKQV KDNENVVNEYSSELEKHQLYIDETVNSNIPTNLR VLRSILENLRSKIQKLESDVSAQMEYCRTPCTVS CNIPVVSGKECEEIIRKGGETSEMYLIQPDSSVKP YRVYCDMNTENGGWTVIQNRQDGSVDFGRKW DPYKQGFGNVATNTDGKNYCGLPGEYWLGNDK ISQLTRMGPTELLIEMEDWKGDKVKAHYGGFTV QNEANKYQISVNKYRGTAGNALMDGASQLMGE NRTMTIHNGMFFSTYDRDNDGWLTSDPRKQCSK EDGGGWWYNRCHAANPNGRYYWGGQYTWDM AKHGTDDGVVWMNWKGSWYSMKKMSMKIRP FFPQQ
3123	A	3	1490	HASGPTRPVSWSFHKLKTMKHLLLLLLCVFLVK SQGVNDNEEGFFSARGHRPLDKKREEAPSLRPAP PPISGGGYRARPAKAAATQKKVERKAPDAGGCL HADPDLGVLCPTGCQLQEALLQQERPIRNSVDEL NNNVEAVSQTSSSSFQYMYLLKDLWQKRQKQV KDNENVVNEYSSELEKHQLYIDETVNSNIPTNLR VLRSILENLRSKIQKLESDVSAQMEYCRTPCTVS CNIPVVSGKECEEIIRKGGETSEMYLIQPDSSVKP YRVYCDMNTENGGWTVIQNRQDGSVDFGRKW DPYKQGFGNVATNTDGKNYCGLPGEYWLGNDK ISQLTRMGPTELLIEMEDWKGDKVKAHYGGFTV QNEANKYQISVNKYRGTAGNALMDGASQLMGE NRTMTIHNGMFFSTYDRDNDGWLTSDPRKQCSK EDGGGWWYNRCHAANPNGRYYWGGQYTWDM AKHGTDDGVVWMNWKGSWYSMKKMSMKIRP FFPQQ
3124	A	3	544	RVDDFVLLRSRLALRWLSHVRRPSRRVPRMPRG SRSRTSRMAPPASRAPQMRAAPRPAPVAQPPAA APPSAVGSSAAAPRQPGLMAQMATTAAGVAVG SAVGHTLGHAITGGFSGGSNAEPARPDITYQEPQ GTQPAQQQQPCLYEIKQFLECAQNQGDIKLCEGF NEVLKQCRLANGLA
3125	A	3	571	GNSYNHRSLAAYPYMSHSQHSPYLQSYHNSSAA AQTRGDDTDQQKTTVIENGEIRFNGKGKKIRKPR TIYSSLQLQALNHRFQQTQYLALPERAELAASLG LTQTQVKIWFQNKRSKFKKLLKQGSNPHESDPL QGSAALSPRSPALPPVWDVSASAKGVSMPPNSY MPGYSHWYSSPHQDTMQRPQMM
3126	A .	43	5377	LSVFFPIPVDGRDRGSNPSLESTSSELSTSTSEGSL SAMSGRNELHSRLHPHPQSSLIPMMFSPPESLLAS CILRGNFAEAHQVLFTFNLKSSPSSGELMFMER Y QEVIQELAQVEHKIENQNSDAGSSTIRRTGSGRST LQAIGSAAAAGMVFYSISDVTDKLLNTSGDPIPM LQEDFWISTALVEPTAPLREVLEDLSPPAMAAFD LACSQCQLWKTCKQLLETAERRLNSSLERRGRRI

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				DHVLLNADGIRGFPVVLQQISKSLNYLLMSASQT KSESVEEKGGGPPRCSITELLQMCWPSLSEDCVA SHTTLSQQLDQVLQSLREALELPEPRTPPLSSLVE QAAQKAPEAEAHPVQIQTQLLQKNLGKQTPSGS RQMDYLGTFFSYCSTLAAVLLQSLSSEPDHVEVK VGNPFVLLQQSSSQLVSHLLFERQVPPERLAALL AQENLSLSVPQVIVSCCCEPLALCSSRQSQQTSSL LTRLGTLAQLHASHCLDDLPLSTPSSPRTTENPTL ERKPYSSPRDSSLPALTSSALAFLKSRSKLLATVA CLGASPRLKVSKPSLSWKELRGRREVPLAAEQV ARECERLLEQFPLFEAFLLAAWEPLRGSLQQGQS LAVNLCGWASLSTVLLGLHSPIALDVLSEAFEES LVARDWSRALQLTEVYGRDVDDLSSIKDAVLSC AVACDKEGWQYLFPVKDASLRSRLALQFVDRW PLESCLEILAYCISDTAVQEGLKCELQRKLAELQ VYQKILGLQSPPVWCDWQTLRSCCVEDPSTVMN MILEAQEYELCEEWGCLYPIPREHLISLHQKHLL HLLERDHDKALQLLRRIPDPTMCLEVTEQSLDQ HTSLATSHFLANYLTTHFYGQLTAVRHREIQALY VGSKILLTLPEQHRASYSHLSSNPLFMLEQLLMN MKVDWATVAVQTLQQLLVGQEIGFTMDEVDSL LSRYAEKALDFPYPQREKRSDSVIHLQEIVHQAA DPETLPRSPSAEFSPAAPPGISSIHSPSLRERSFPPT QPSQEFVPPATPPARHQWVPDETESICMVCCREH FTMFNRRHHCRRCGRLVCSSCSTKKMVVEGCRE NPARVCDQCYSYCNKDVPEEPSEKPEALDSSKSE SPPYSFVVRVPKADEVEWILDLKEEENELVRSEF YYEQAPSASLCIAILNLHRDSIACGHQLIEHCCRL SKGLTNPEVDAGLLTDIMKQLLFSAKMMFVKAG QSQDLALCDSYISKVDVLNILVAAAYRHVPSLDQ ILQPAAVTRLRNQLLEAEYYQLGVEVSTKTGLDT TGAWHAWGMACLKAGNLTAAREKFSRCLKPPF DLNQLNHGSRLVQDVVEYLESTVRPFVSLQDDD YFATLRELEATLRTQSLSLAVIPEGKIMNNTYYQ ECLFYLHNYSTNLAIISFYVRHSCLREALLHLLNK ESPPEVFIEGIFQPSYKSGKLHTLENLLESIDPTLES WGKYLIAACQHLQKKNYYHILYELQQFMKDQV RAAMTCIRFFSHKAKSYTELGEKLSWLLKAKDH LKIYLQETSRSSGRKKTTFFRKKMTAADVSRHM NTLQLMEVTRFLHRCESAGTSQITTLPLPTLFG NNHMKMDVACKVMLGGKNVEDGFGIAFRVLQ DFQLDAAMTYCRAARQLVEKEKYSEIQQLLKCV SESGMAAKSDGDTILLNCLEAFKRIPPQCCFCSA QELEGLIQAIHNDDNKVRAYLICCKLRSAYLIAV KQEHSRATALVQQVQQAAKSSGDAVVQDICAQ
3127	A	467	1259	WLLTSHPRGAHGPGSRK HLGPPLAWIPAASLTSTKGEFGVEDDRPARGPPP PKSEEASWSESGVSSSSGDGPFAGGEVDKRLHQL KTQLATLTSSLATVTQEKSRMEASYLADKKKMK QDLEDASNKAEEERARLEGELKGLQEQIAETKA RLITQQHDRAQEQSDHALMLRELQKLLQEERTQ RQDLELRLEETREALAGRAYAAEQMEGFELQTK
3128	A	1854	798	QLTREVEELKSELQAIRDEKNQPDPRLQELQEEA ARLKSHFQAQLQQEMRKVIIHISFKHQPLT ASGSPAPSSSSAMAAACGPGAAGYCLLLGLHLFL

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				LTAGPALGWNDPDRMLLRDVKALTLHYDRYTT SRRLDPIPQLKCVGGTAGCDSYTPKVIQCQNKG WDGYDVQWECKTDLDIAYKFGKTVVSCEGYES SEDQYVLRGSCGLEYNLDYTELGLQKLKESGKQ HGFASFSDYYYKWSSADSCNMSGLITIVVLLGIA FVVYKLFLSDGQYSPPPYSEYPPFSHRYQRFTNS AGPPPPGFKSEFTGPQNTGHGATSGFGSAFTGQQ GYENSGPGFWTGLGTGGILGYLFGSNRAATPFSD SWYYPSYPPSYPGTWNRAYSPLHGGSGSYSVCS NSDTKTRTASGYGGTRRR
3129	A	2340	1192	ELARPKQQSSEKSRNMIRNWLTIFILFPLKLVEK CESSVSLTVPPVVKLENGSSTNVSLTLRPPLNATL VITFEITFRSKNITILELPDEVVVPPGVTNSSFQVT SQNVGQLTVYLHGNHSNQTGPRIRFLVIRSSAISII NQVIGWIYFVAWSISFYPQVIMNWRRKSVIGLSF DFVALNLTGFVAYSVFNIGLLWVPYIKEQFLLKY PNGVNPVNSNDVFFSLHAVVLTLIIIVQCCLYERG GQRVSWPAIGFLVLAWLFAFVTMIVAAVGVITW LQFLFCFSYIKLAVTLVKYFPQAYMNFYYKSTEG WSIGNVLLDFTGGSFSLLQMFLQSYNNDQWTLIF GDPTKFGLGVFSIVFDVVFFIQHFCLYRKRPGYD QLN
3130	A	31	2026	CWWPPLLPQLEPEPPPLRPRVAASQGGGMLGKG VVGGGGGTKAPKPSFVSYVRPEEIHTNEKEVTEK EVTLHLLPGEQLLCEASTVLKYVQEDSCQHGVY GRLVCTDFKIAFLGDDESALDNDETQFKNKVIGE NDITLHCVDQIYGVFDEKKKTLFGQLKKYPEKLII HCKDLRVFQFCLRYTKEEEVKRIVSGIIHHTQAP KLLKRLFLFSYATAAQNNTVTDPKNHTVMFDTL KDWCWELERTKGNMKYKAVSVNEGYKVCERL PAYFVVPTPLPEENVQRFQGHGIPIWCWSCHNGS ALLKMSALPKEQDDGILQIQKSFLDGIYKTIHRPP YEIVKTEDLSSNFLSLQEIQTAYSKFKQLFLIDNST EFWDTDIKWFSLLESSSWLDIIRRCLKKAIEITEC MEAQNMNVLLLEENASDLCCLISSLVQLMMDPH CRTRIGFQSLIQKEWVMGGHCFLDRCNHLRQND KEEHQRQLSLPLTQSKSSPKRGFFREETDHLIKNL LGKRISKLINSSDELQDNFREFYDSWHSKSTDYH GLLLPHIEGPEIKVWAQRYLRWIPEAQILGGGQV ATLSKLLEMMEEVQSLQEKIDERHHSQQAPQAE APCLLRNSARLSSLFPFALLQRHSSKPVLPTSGW KALGDEDDLAKREDEFVDLGDV
3131	A	126	965	QSRSRPRREGVGTGSRAVLCILATCGSKMSDIGD WFRSIPAITRYWFAATVAVPLVGKLGLISPAYLF LWPEAFLYRFQIWRPITATFYFPVGPGTGFLYLV NLYFLYQYSTRLETGAFDGRPADYLFMLLFNWI CIVITGLAMDMQLLMIPLIMSVLYVWAQLNRDM IVSFWFGTRFKACYLPWVILGFNYIIGGSVINELIG NLVGHLYFFLMFRYPMDLGGRNFLSTPQFLYRW LPSRRGGVSGFGVPPASMRRAADQNGGGGRHN WGQGFRLGDQ
3132	A	2	350	FVAGWRALTAPSTSARLRAFGWQAAARLLVFG ARGVGLGSGAPGSLPCYLRMDALALLGGLVNV ARLPERWGPGRFDYWGNSHQIMHLLSVGSILQL HAGVVPDLLWAAHHACPRD

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3133	A	1	2921	MTCFKGQKGEQRSHAFEANKDHKAKVPSPNLYS QLNALQFTVDERSILWLNQFLLDLKQSLNQFMA VYKLNDNSKSDEHVDVRVDGLMLKFVIPSEVKS ECHQDQPRAISIQSSEMIATNTRHCPNCRHSDLEA LFQDFKDCDFFSKTYTSFPKSCDNFNLLHPIFQRH AHEQDTKMHEIYKGNITPQLNKNTLKTSAATDV WAVYFSQFWIDYEGMKSGKGRPISFVDSFPLSIW ICQPTRYAESQKEPQTCNQVSLNTSQSESSDLAG RLKRKKLLKEYYSTESEPLTNGGQKPSSSDTFFR FSPSSSEADIHLLVHVHKHVSMQINHYQYLLLLF LHESLILLSENLRKDVEAVTGSPASQTSICIGILLR SAELALLLHPVDQANTLKSPVSESVSPVVPDYLP TENGDFLSSKRKQISRDINRIRSVTVNHMSDNRS MSVDLSHIPLKDPLLFKSASDTNLQKGISFMDYL SDKHLGKISEDESSGLVYKSGSGEIGSETSDKKDS FYTDSSSVLNYREDSNILSFDSDGNQNILSSTLTS KGNETIESIFKAEDLLPEAASLSENLDISKEETPPV RTLKSQSSLSGKPKERCPPNLAPLCVSYKNMKRS SSQMSLDTISLDSMILEEQLLESDGSDSHMFLEKG NKKNSTTNYRGTAESVNAGANLQNYGETSPDAI STNSEGAQENHDDLMSVVVFKITGVNGEIDIRGE DTEICLQVNQVTPDQLGNISLRHYLCNRPVGSDQ KAVIHSKSSPEISLRFESGPGAVIHSLLAEKNGFL
				QCHIENFSTEFLTSSLMNIQHFLEDETVATVMPM KIQVSNTKINLKDDSPRSSTVSLEPAPVTVHIDHL VVERSDDGSFHIRDSHMLNTGNDLKENVKSDSV LLTSGKYDLKKQRSVTQATQTSPGVPWPSQSAN FPEFSFDFTREQLMEENESLKQELAKAKMALAE AHLEKDALLHHIKKMTVE
3134		9	1579	EEEGLSGGGPRVPCSLWGKQTMDYDFKAKLAA ERERVEDLFEYEGCKVGRGTYGHVYKARRKDG KDEKEYALKQIEGTGISMSACREIALLRELKHPN VIALQKVFLSHSDRKVWLLFDYAEHDLWHIIKFH RASKANKKPMQLPRSMVKSLLYQILDGIHYLHA NWVLHRDLKPANILVMGEGPERGRVKIADMGF ARLFNSPLKPLADLDPVVVTFWYRAPELLLGAR HYTKAIDIWAIGCIFAELLTSEPIFHCRQEDIKTSN PFHHDQLDRIFSVMGFPADKDWEDIRKMPEYPT LQKDFRRTTYANSSLIKYMEKHKVKPDSKVFLL LQKLLTMDPTKRITSEQALQDPYFQEDPLPTLDV FAGCQIPYPKREFLNEDDPEEKGDKNQQQQNQ HQQPTAPPQQAAAPPQAPPPQQNSTQTNGTAGG AGAGVGGTGAGLQHSQDSSLNQVPPNKKPRLGP SGANSGGPVMPSDYQHSSSRLNYQSSVQGSSQS QSTLGYSSSSQQSSQYHPSHQAHRY
3135	Α .	3		ERKMAEPPSPVHCVAAAAPTATVSEKEPFGKLQ LSSRDPPGSLSAKKVRTEEKKAPRRVNGEGGSG GNSRQLQPPAAPSPQSYGSPASWSFAPLSAAPSPS SSRSSFSFSAGTAVPSSASASLSQPGPRKLLVPPTL LHAQPHHLLLPAAAAAASANAKSRRPKEKREKE RRRHGLGGAREAGGASREENGEVKPLPRDKIKD KIKERDKEKEREKKKHKVMNEIKKENGEVKILL KSGKEKPKTNIEDLQIKKVKKKKKKKHKENEKR KRPKMYSKSIQTICSGLLTDVEDQAAKGILNDNI KDYVGKNLDTKNYDSKIPENSEFPFVSLKEPRVQ

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SEQ ID NO:	Method	Predicted beginning nucleotide location corresponding to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion
	ļ			NNLKRLDTLEFKQLIHIEHQPNGGASVIHCLQ
3136	A	1442	682	TAAMSIFTPTNQIRLTNVAVVRMKRAGKRFEIAC YKNKVVGWRSGVEKDLDEVLQTHSVFVNVSKG QVAKKEDLISAFGTDDQTEICKQILTKGEVQVSD KERHTQLEQMFRDIATIVADKCVNPETKRPYTVI LIERAMKDIHYSVKTNKSTKQQALEVIKQLKEK MKIERAHMRLRFILPVNEGKKLKEKLKPLIKVIES EDYGQQLEIVCLIDPGCFREIDELIKKETKGKGSL EVLNLKDVEEGDEKFE
3137	A		3143	MVEGKRHVLHGGRQERMRAKQKGKPLIKSSDL VRLIHYHHNSSPLHKQSSGPSSSPAAAAAPEKPG PKAAEVGDDFLGDFVVGERVWVNGVKPGVVQY LGETQFAPGQWAGVVLDDPVGKNDGAVGGVR YFECPALQGIFTRPSKLTRQPTAEGSGSDAHSVES LTAQNLSLHSGTATPPLTSRVIPLRESVLNSSVKT GNESGSNLSDSGSVKRGEKDLRLGDRVLVGGTK TGVVRYVGETDFAKGEWCGVELDEPLGKNDGA VAGTRYFQCPPKFGLFAPIHKVIRIGFPSTSPAKA KKTKRMAMGVSALTHSPSSSSISSVSSVASSVGG RPSRSGLLTETSSRYARKISGTTALQEALKEKQQ HIEQLLAERDLERAEVAKATSHICEVEKEIALLK AQHEQYVAEAEEKLQRARLLVESVRKEKVDLSN QLEEERRKVEDLQFRVEEESITKGDLETQTQLEH ARIGELEQSLLLEKAQAERLLRELADNRLTTVAE KSRVLQLEEELTLRRGEIEELQQCLLHSGPPPPDH PDAAEILRLRERLLSASKEHQRESGVLRDKYEKA LKAYQAEVDKLRAANEKYAQEVAGLKDKVQQ ATSENMGLMDNWKSKLDSLASDHQKSLEDLKA TLNSGPGAQQKEIGELKAVMEGIKMEHQLELGN LQAKHDLETAMHVKEKEALREKLQEAQEELAG LQRHWRAQLEVQASQHRLELQEAQDQRRDAEL RVHELEKLDVEYRGQAQAIEFLKEQISLAEKKML DYERLQRAEAQGKQEVESLREKLLVAENRLQAV EALCSSQHTHMIESNDISEETIRTKETVEGLQDKL NKRDKEVTALTSQTEMLRAQVSALESKCKSGEK KVDALLKEKRRLEAELETVSRKTHDASGQLVLIS QELLRKERSLNELRVLLLEANRHSPGPERDLSRE VHKAEWRIKEQKLKDDIRGLREKLTGLDKEKSL SDQRRYSLIDPSSAPELLRLQHQLMSTEDALRDA LDQAQQVEKLMEAMRSCPDKAQTIGNSGSANGI HQQDKAQKQEDKH
3138	A	110	2499	QDRRLLRLELQKTCQPTSTMSGSHTPACGPFSAL TPSIWPQEILAKYTQKEESAEQPEFYYDEFGFRV YKEEGDEPGSSLLANSPLMEDAPQRLRWQAHLE FTHNHDVGDLTWDKIAVSLPRSEKLRSLVLAGIP HGMRPQLWMRLSGALQKKRNSELSYREIVKNSS NDETIAAKQIEKDLLRTMPSNACFASMGSIGVPR LRRVLRALAWLYPEIGYCQGTGMVAACLLLFLE EEDAFWMMSAIIEDLLPASYFSTTLLGVQTDQRV LRHLIVQYLPRLDKLLQEHDIELSLITLHWFLTAF ASVVDIKLLLRIWDLFFYEGSRVLFQLTLGMLHL KEEELIQSENSASIFNTLSDIPSQMEDAELLLGVA MRLAGSLTDVAVETQRRKHLAYLIADQGQLLGA GTLTNLSQVVRRRTQRRKSTITALLFGEDDLEAL KAKNIKQTELVADLREAILRVARHFQCTDPKNCS

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Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, SEO ID Method Predicted Predicted end E-Glutamic Acid, F-Phenylalanine, G-Glycine, H-Histidine, nucleotide NO: beginning I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, nucleotide location N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, corresponding location corresponding T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, to last amino X=Unknown, *=Stop codon, /=possible nucleotide deletion, acid residue of to first amino acid residue of \=possible nucleotide insertion peptide peptide sequence sequence VVSRQLPGLLPNTALTPPTPLVGLCSLWQELTPD YSMESHQRDHENYVACSRSHRRRAKALLDFERH DDDELGFRKNDIITIVSQKDEHCWVGELNGLRG WFPAKFVEVLDERSKEYSIAGDDSVTEGVTDLV RGTLCPALKALFEHGLKKPSLLGGACHPWLFIEE AAGREVERDFASVYSRLVLCKTFRLDEDGKVLT PEELLYRAVOSVNVTHDAVHAQMDVKLRSLICV GLNEOVLHLWLEVLCSSLPTVEKWYQPWSFLRS PGWVOIKCELRVLCCFAFSLSQDWELPAKREAQ **OPLKEGVRDMLVKHHLFSWDVDG** 3139 110 2499 ODRRLLRLELOKTCOPTSTMSGSHTPACGPFSAL Α TPSIWPQEILAKYTQKEESAEQPEFYYDEFGFRV YKEEGDEPGSSLLANSPLMEDAPQRLRWQAHLE FTHNHDVGDLTWDKIAVSLPRSEKLRSLVLAGIP HGMRPQLWMRLSGALQKKRNSELSYREIVKNSS NDETIAAKQIEKDLLRTMPSNACFASMGSIGVPR LRRVLRALAWLYPEIGYCQGTGMVAACLLLFLE **EEDAFWMMSAIIEDLLPASYFSTTLLGVQTDQRV** LRHLIVQYLPRLDKLLQEHDIELSLITLHWFLTAF ASVVDIKLLLRIWDLFFYEGSRVLFQLTLGMLHL KEEELIQSENSASIFNTLSDIPSQMEDAELLLGVA MRLAGSLTDVAVETQRRKHLAYLIADQGQLLGA GTLTNLSQVVRRRTQRRKSTITALLFGEDDLEAL KAKNIKQTELVADLREAILRVARHFQCTDPKNCS VVSRQLPGLLPNTALTPPTPLVGLCSLWQELTPD YSMESHORDHENYVACSRSHRRRAKALLDFERH DDDELGFRKNDIITIVSQKDEHCWVGELNGLRG WFPAKFVEVLDERSKEYSIAGDDSVTEGVTDLV RGTLCPALKALFEHGLKKPSLLGGACHPWLFIEE AAGREVERDFASVYSRLVLCKTFRLDEDGKVLT PEELLYRAVQSVNVTHDAVHAQMDVKLRSLICV GLNEOVLHLWLEVLCSSLPTVEKWYQPWSFLRS PGWVQIKCELRVLCCFAFSLSQDWELPAKREAQ OPLKEGVRDMLVKHHLFSWDVDG 4939 SAALGASLAIPRPGLPGVHGRGPGTLSGRAMEG 3140 1 **AEPRARPERLAEAETRAADGGRLVEVQLSGGAP** WGFTLKGGREHGEPLVITKIEEGSKAAAVDKLL **AGDEIVGINDIGLSGFRQEAICLVKGSHKTLKLV** VKRRSELGWRPHSWHATKFSDSHPELAASPFTST SGCPSWSGRHHASSSSHDLSSSWEQTNLQRTLD HFSSLGSVDSLDHPSSRLSVAKSNSSIDHLGSHSK RDSAYGSFSTSSSTPDHTLSKADTSSAENILYTVG LWEAPROGGROAOAAGDPQGSEEKLSCFPPRVP GDSGKGPRPEYNAEPKLAAPGRSNFGPVWYVPD KKKAPSSPPPPPPPLRSDSFAATKSHEKAQGPVFS EAAAAQHFTALAQAQPRGDRRPELTDRPWRSAH **PGSLGKGSGGPGCPQEAHADGSWPPSKDGASSR** LQASLSSSDVRFPQSPHSGRHPPLYSDHSPLCADS LGOEPGAASFONDSPPOVRGLSSCDOKLGSGWQ **GPRPCVQGDLQAAQLWAGCWPSDTALGALESL PPPTVGQSPRHHLPQPEGPPDARETGRCYPLDKG AEGCSAGAQEPPRASRAEKASQRLAASITWADG ESSRICPQETPLLHSLTQEGKRRPESSPEDSATRPP** PFDAHVGKPTRRSDRFATTLRNEIQMHRAKLQK SRSTVALTAAGEAEDGTGRWRAGLGGGTQEGPL O VAIO. AVV

SEQ ID NO:	Method	Predicted beginning nucleotide location corresponding to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isòleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion
				AGTYKDHLKEAQARVLRATSFKRRDLDPNPGDL YPESLEHRMGDPDTVPHFWEAGLAQPPSSTSGGP HPPRIGGRRRFTAEQKLKSYSEPEKMNEVGLTRG YSPHQHPRTSEDTVGTFADRWKFFEETSKPVPQR PAQKQALHGIPRDKPERPRTAGRTCEGTEPWSRT TSLGDSLNAHSAAEKAGTSDLPRRLGTFAEYQAS WKEQRKPLEARSSGRCHSADDILDVSLDPQERPQ HVHGRSRSSPSTDHYKQEASVELRRQAGDPGEP REELPSAVRAEEGQSTPRQADAQCREGSPGSQQ HPPSQKAPNPPTFSELSHCRGAPELPREGRGRAG TLPRDYRYSEESTPADLGPRAQSPGSPLHARGQD SWPVSSALLSKRPAPQRPPPPKREPRRYRATDGA PADAPVGVLGRPFPTPSPASLDVYVARLSLSHSPS VFSSAQPQDTPKATVCERGSQHVSGDASRPLPEA LLPPKQQHLRLQTATMETSRSPSPQFAPQKLTDK PPLLIQDEDSTRIERVMDNNTTVKMVPIKIVHSES QPEKESRQSLACPAEPPALPHGLEKDQIKTLSTSE QFYSRFCLYTRQGAEPEAPHRAQPAEPQPLGTQV PPEKDRCTSPPGLSYMKAKEKTVEDLKSEELARE IVGKDKSLADILDPSVKIKTTMDLMEGIFPKDEH LLEEAQQRRKLLPKIPSPRSTEERKEEPSVPAAVS LATNSTYYSTSAPKAELLIKMKDLQEQQEHEEDS GSDLDHDLSVKKQELIESISRKLQVLREARESLLE DVQANTVLGAEVEAIVKGVCKPSEFDKFRMFIG
•				DLDKVVNLLLSLSGRLARVENALNNLDDGASPG DRQSLLEKQRVLIQQHEDAKELKENLDRRERIVF DILANYLSEESLADYEHFVKMKSALIIEQRELED KIHLGEEQLKCLLDSLQPERGK
3141	A	97	1894	SPRGATMETPPLPPACTKQGHQKPLDSKDDNTE KHCPVTVNPWHMKKAFKVMNELRSQNLLCDVT IVAEDMEISAHRVVLAACSPYFHAMFTGEMSESR AKRVRIKEVDGWTLRMLIDYVYTAEIQVTEENV QVLLPAAGLLQLQDVKKTCCEFLESQLHPVNCL GIRAFADMHACTDLLNKANTYAEQHFADVVLSE EFLNLGIEQVCSLISSDKLTISSEEKVFEAVIAWV NHDKDVRQEFMARLMEHVRLPLLPREYLVQRV EEEALVKNSSACKNYLIEAMKYHLLPTEQRILMK SVRTRLRTPMNLPKLMVVVGGQAPKAIRSAECY DFKEQRWHQVAELPSRRCRAGMVYLAGLVFAV GGFNGSLRVRTVDSYDPVKDQWTSVANMRDRR STLGAAVLNGLLYAVGGFDGSTGLSSVEAYNIKS NEWFHVAPMNTRRSSVGVGVVGGLLYAVGGYD GASRQYLSTVECYNATTNEWTYIAEMSTRRSGA GVGVLNNLLYAVGGHDGPLVRKSVEVYDPTTN AWRQVADMNMCRRNAGVCAVNGLLYVVGGD DGSCNLASVEYYNPTTDKWTVVSSCMSTGRSYA GVTVIDKPL
3142	A	1211	1311	FSNLTTEKVAHAKEENLSMHQMLDQTLLELNN M
3143	A	1809	1041	SEELDREKKLKEDSPRKTPNKESGVPSLPVSLTSI KEEPKEAKHPDSQSMEESKLKNDDRKTPVNWK DSRGTRVAVSSPMSQHQSYIQYLHAYPYPQMYD PSHPAYRAVSPVLMHSYPGAYLSPGFHYPVYGK MSGREETEKVNTSPSVNTKTTTESKALDLLQQH ANQYRSKSPAPVEKATAEREREAERERDRHSPFG

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				QRHLHTHHHTHVGMGYPLIPGQYDPFQGLTSAA LVASQQVAAQASASGMFPGQRR
3144	A	78	604	SVSGIVLDLLPYLHFLSNMNLDGSAQDPEKREYS SVCVGREDDIKKSERMTAVVHDREVVIFYHKGE YHAMDIRCYHSGGPLHLGDIEDFDGRPCIVCPW HKYKITLATGEGLYQSINPKDPSAKPKWCSKGIK QRIHTVTVDNGNIYVTLSNEPFKCDSDFYATGDF KVIKSSS
3145	A	2	333	RNSLLLPPLHLDNSTPAKMSCQQNQQQCQPPPK CPSPKCPPKSPVQCLPPASSGCAPSSGGCGPSSEG GCFLNHHRRHHRCRRQRPNSCDRGSGQQGGGS GCGHGSGGCC
3146	A	3	1151	VCTALQEFGTRSTLLRCLDSGFRPGASRGLVGSW AAMESTLGAGIVIAEALQNQLAWLENVWLWITF LGDPKILFLFYFPAAYYASRRVGIAVLWISLITEW LNLIFKWFLFGDRPFWWVHESGYYSQAPAQVHQ FPSSCETGPGSPSGHCMITGAALWPIMTALSSQV ATRARSRWVRVMPSLAYCTFLLAVGLSRIFILAH FPHQVLAGLITGAVLGWLMTPRVPMERELSFYG LTALALMLGTSLIYWTLFTLGLDLSWSISLAFKW CERPEWIHVDSRPFASLSRDSGAALGLGIALHSPC YAQVRRAQLGNGQKIACLVLAMGLLGPLDWLG HPPQISLFYIFNFLKYTLWPCLVLALVPWAVHMF SAQEAPPIHSS
3147	A	1437	594	RSFSLSFSLLSPSEMMALGAAGATRVFVAMVAA ALGGHPLLGVSATLNSVLNSNAIKNLPPPLGGAA GHPGSAVSAAPGILYPGGNKYQTIDNYQPYPCAE DEECGTDEYCASPTRGGDAGVQICLACRKRKR CMRHAMCCPGNYCKNGICVSSDQNHFRGEIEETI TESFGNDHSTLDGYSRRTTLSSKMYHTKGQEGS VCLRSSDCASGLCCARHFWSKICKPVLKEGQVC TKHRRKGSHGLEIFQRCYCGEGLSCRIQKDHHQ ASNSSRLHTCQRH
3148	A		1562	MSTLYDIRAHKAQLLRFFASSDSNKALEQRRTLH TPKLEHLDRVLYEWFLGKRSEGVPVSGPMLIEK AKDFYEQMQLTEPCVFSGGWLWRFKARHGIKK LDASSEKQSADHQAAEQFCAFFRSLAAEHGLSA EQVYNADETGLFWRCLPNPTPEGGAVPGPKQGK DRLTVLMCANATGSHRLKPLAIGKCSGPRAFKGI QHLPVAYKAQGNAWVDKEIFSDWFHHIFVPSVR EHFRTIGLPEDSKAVLLLDSSRAHPQEAELVSSN VFTIFLPASVASLVQPMEQGIRRDFMRNFINPPVP LQGPHARYNMNDAIFSVACAWNAVPSHVFRRA WRKLWPSVAFAEGSSSEEELEAECFPVKPHNKSF AHILELVKEGSSCPGQLRQRQAASWGVAGREAE GGRPPAATSPAEVVWSSEKTPKADQDGRGDPGE GEEVAWEQAAVAFDAVLRFAERQPCFSAQEVG QLRALRAVFRSQQQVRRRRGALGAVVKVEALQ EGPGGCGATAQSPLPCSSTAGDN
3149	A	132	4125	VAVMISTAPLYSGVHNWTSSDRIRMCGINEERRA PLSDEESTTGDCQHFGSQEFCVSSSFSKVELTAV GSGSNARGADPDGSATEKLGHKSEDKPDDPQPK MDYAGNVAEAEGLLVPLSSPGDGLKLPASDSAE ASNSRADCSWTPLNTQMSKQVDCSPAGVKALDS RQGVGEKNTFILATLGTGVPVEGTLPLVTTNFSP

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SEQ ID NO:	Method	Predicted beginning nucleotide location corresponding to first amino acid residue of peptide	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion
		sequence		LPAPICPPAPSSASVPHSVPDAFQAPVPPSAPTLVL APVPTPVLAPMPASTPPAAPAPPSVPMPTPTPSSG PPSTPTLIPAFAPTPVPAPTPAPIFTPAPTPMPAATP AAIPTSAPIPASFSLSRVCFPAAQAPAMQKVPLSF QPGTVLTPSQPLVYIPPPSCGQPLSVATLPTTLGV SSTLTLPVLPSYLQDRCLPGVLASPELRSYPYAFS VARPLTSDSKLVSLEVNRLPCTSPSGSTTTQPAPD GVPGPLADTSLVTASAKVLPTPQPLLPAPSGSSAP PHPAKMPSGTEQQTEGTSVTFSPLKSPPQLEREM ASPPECSEMPLDLSSKSNRQKLPLPNQRKTPPMP VLTPVHTSSKALLSTVLSRSQRTTQAAGGNVTSC LGSTSSPFVIFPEIVRNGDPSTWVKNSTALISTIPG TYVGVANPVPASLLLNKDPNLGLNRDPRHLPKQ EPISIIDQGEPKGTGATCGKKGSQAGAEGQPSTV KRYTPARIAPGLPGCQTKELSLWKPTGPANIYPR CSVNGKPTSTQVLPVGWSPYHQASLLSIGISSAG QLTPSQGAPIRPTSVVSEFSGVPSLSSSEAVHGLP EGQPRPGGSFVPEQDPVTKNKTCRIAAKPYEEQV NPVLLTLSPQTGTLALSVQPSGGDIRMNQGPEES ESHLCSDSTPKMEGPQGACGLKLAGDTKPKNQV LATYMSHELVLATPQNLPKMPELPLLPHDSHPKE LILDVVPSSRRGSSTERPQLGSQVDLGRVKMEKV DGDVVFNLATCFRADGLPVAPQRGQAEVRAKA GQARVKQESVGVFACKNKWQPDDVTESLPPKK MKCGKEKDSEEQQLQPQAKAVVRSSHRPKCRK LPSDPQESTKKSPRGASDSGKEHNGVRGKHKHR KPTKPESQSPGKRADSHEEGSLEKKAKSSFRDFIP VVLSTRTRSQSDLKARKQKTSSSQSLEHRLRNRN LLLPNKVQGISDSPNGFLPNNLEEPACLENSEKPS GKRKCKTKHMATVSEEAKGKGRWSQQKTRSPK SPTPVKPTEPCTPSKSRSASSEEASESPTARQIPPE ARRLIVNKNAGETLLQRAARLGYKDVVLYCLQK DSEDVNHRDNAGYTALHEACSRGWTDILNILLE
3150	A	3	2795	SLRMHNLSILVRQIKFYYQETLQQLIMMSLPNVLI IGKNPFSEQGTEEVKKLLLLLLGCAVQCQKKEEF IERIQGLDFDTKAAVAAHIQEVTHNQENVFDLQ WMEVTDMSQEDIEPLLKNMALHLKRLIDERDEH SETIIELSEERDGLHFLPHASSSAQSPCGSPGMKR TESRQHLSVELADAKAKIRRLRQELEEKTEQLLD CKQELEQMEIELKRLQQENMNLLSDARSARMYR DELDALREKAVRVDKLESEVSRYKERLHDIEFY KARVEELKEDNQVLLETKTMLEDQLEGTRARSD KLHELEKENLQLKAKLHDMEMERDMDRKKIEE LMEENMTLEMAQKQSMDESLHLGWELEQISRTS ELSEAPQKSLGHEVNELTSSRLLKLEMENQSLTK TVEELRTTVDSVEGNASKILKMEKENQRLSKKV EILENEIVQEKQSLQNCQNLSKDLMKEKAQLEKT IETLRENSERQIKILEQENEHLNQTVSSLRQRSQIS AEARVKDIEKENKILHESIKETSSKLSKIEFEKRQI KKELEHYKEKGERAEELENELHHLEKENELLQK KITNLKITCEKIEALEQENSELERENRKLKKTLDS FKNLTFQLESLEKENSQLDEENLELRRNVESLKC ASMKMAQLQLENKELESEKEQLKKGLELLKASF KKTERLEVSYQGLDIENQRLQKTLENSNKKIQQL

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				ESELQDLEMENQTLQKNLEELKISSKRLEQLEKE NKSLEQETSQLEKDKKQLEKENKRLRQQAEIKD TTLEENNVKIGNLEKENKTLSKEIGIYKESCVRLE ELEKENKELVKRATIDIKTLVTLREDLVSEKLKT QQMNNDLEKLTHELEKIGLNKERLLHDEQSTDD SRYKLLESKLESTLKKSLEIKEEKIAALEARLEES TNYNQQLRQELKTVKKK
3151	A	2	2515	GFWLHLTLLGASLPAALGWMDPGTSRGPDVGV GESQAEEPRSFEVTRREGLSSHNELLASCGKKFC SRGSRCVLSRKTGEPECQCLEACRPSYVPVCGSD GRFYENHCKLHRAACLLGKRITVIHSKDCFLKGD TCTMAGYARLKNVLLALQTRLQPLQEGDSRQDP ASQKRLLVESLFRDLDADGNGHLSSSELAQHVL KKQDLDEDLLGCSPGDLLRFDDYNSDSSLTLREF YMAFQVVQLSLAPEDRVSVTTVTVGLSTVLTCA VHGDLRPPIIWKRNGLTLNFLDLEDINDFGEDDS LYITKVTTIHMGNYTCHASGHEQLFQTHVLQVN VPPVIRVYPESQAQEPGVAASLRCHAEGIPMPRIT WLKNGVDVSTQMSKQLSLLANGSELHISSVRYE DTGAYTCIAKNEVGVDEDISSLFIEDSARKTLANI LWREEGLSVGNMFYVFSDDGIIVIHPVDCEIQRH LKPTEKIFMSYEEICPQREKNATQPCQWVSAVNV RNRYIYVAQPALSRVLVVDIQAHKVLQSIGVDPL PAKLSYDKSHDQVWVLSWGDVHKSRPSLQVITE ASTGQSQHLIRTPFAGVDDFFIPPTNLIINHIRFGFI FNKSDPAVHKVDLETMMPLKTIGLHHHGCVPQA MAHTHLGGYFFIQCRQDSPASAARQLLVDSVTD SVLGPNGDVTGTPHTSPDGRFIVSAAADSPWLHV QEITVRGEIQTLYDLQINSGISDLAFQRSFTESNQ YNIYAALHTEPDLLFLELSTGKVGMLKNLKEPPA GPAQPWGGTHRIMRDSGLFGQYLLTPARESLFLI NGRQNTLRCEVSGIKGGTTVVWVGEV
3152	A		2645	GAGWQVSLTGRWSPGREAGAGEVRQDPGSTAA SPSSCDADLSARMARGERRRAVPAEGVRTAER AARGGPGRRDGRGGGPRSTAGGVALAVVVLSL ALGMSGRWVLAWYRARRAVTLHSAPAVLPADS SSPAVAPDLFWGTYRPHVYFGMKTRSPKPLLTG LMWAQQGTTPGTPKLRHTCEQGDGVGPYGWEF HDGLSFGRQHIQDGALRLTTEFVKRPGGQHGGD WSWRVTVEPQDSGTSALPLVSLFFYVVTDGKEV LLPEVGAKGQLKFISGHTSELGDFRFTLLPPTSPG DTAPKYGSYNVFWTSNPGLPLLTEMVKSRLNSW FQHRPPGASPERYLGLPGSLKWEDRGPSGQGQG QFLIQQVTLKIPISIEFVFESGSAQAGGNQALPRLA GSLLTQALESHAEGFRERFEKTFQLKEKGLSSGE QVLGQAALSGLLGGIGYFYGQGLVLPDIGVEGSE QKVDPALFPPVPLFTAVPSRSFFPRGFLWDEGFH QLVVQRWDPSLTREALGHWLGLLNADGWIGRE QILGDEARARVPPEFLVQRAVHANPPTLLLPVAH MLEVGDPDDLAFLRKALPRLHAWFSWLHQSQA GPLPLSYRWRGRDPALPTLLNPKTLPSGLDDYPR ASHPSVTERHLDLRCWVALGARVLTRLAEHLGE AEVAAELGPLAASLEAAESLDELHWAPELGVFA DFGNHTKAVQLKPRPPQGLVRVVGRPQPQLQYV DALGYVSLFPLLLRLLDPTSSRLGPLLDILADSRH

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SEQ ID NO:	Method	Predicted beginning nucleotide location corresponding to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion
				LWSPFGLRSLAASSSFYGQRNSEHDPPYWRGAV WLNVNYLALGALHHYGHLEGPHQARAAKLHGE LRANVVGNVWRQYQATGFLWEQYSDRDGRGM GCRPFHGWTSLVLLAMAEDY
3153	A		4312	MVIKTDELPAAAPADSAREHGSQAGGKGRPGAA AVLLADLERDARQGECALPGAAMAGLAPLKPE ASRSSSPGPTGCIRARVAAEAGTRNPGNAGAELE SWLPCCHGHPETPEPRGGQLPTAPELPSVMLLNG DCPESLKKEAAAAEPPRENGLDEAGPGDETTGQ EVIVIQDTGFSVKILAPGIEPFSLQVSPQEMVQEIH QVLMDREDTCHRTCFSLHLDGNVLDHFSELRSV EGLQEGSVLRVVEEPYTVREARIHVRHVRDLLKS LDPSDAFNGVDCNSLSFLSVFTDGDLGDSGKRK KGLEMDPIDCTPPEYILPGSRERPLCPLQPQNRD WKPLQCLKVLTMSGWNPPPGNRKMHGDLMYLF VITAEDRQVSITASTRGFYLNQSTAYHFNPKPASP RFLSHSLVELLNQISPTFKKNFAVLQKKRVQRHP FERIATPFQVYSWTAPQAEHAMDCVRAEDAYTS RLGYEEHIPGQTRDWNEELQTTRELPRKNLPERL LRERAIFKVHSDFTAAATRGAMAVIDGNVMAIN PSEETKMQMFIWNNIFFSLGFDVRDHYKDFGGD VAAYVAPTNDLNGVRTYNAVDVEGLYTLGTVV VDYRGYRVTAQSIIPGILERDQEQSVIYGSIDFGK TVVSHPRYLELLERTSRPLKILRHQVLNDRDEEV ELCSSVECKGIIGNDGRHYILDLLRTFPPDLNFLP VPGEELPEECARAGFPRAHRHKLCCLRQELVDA FVEHRYLLFMKLAALQLMQQNASQLETPSSLEN GGPSSLESKSEDPPGQEAGSEEGSSASGLAKVK ELAETIAADDGTDPRSREVIRNACKAVGSISSTAF DIRFNPDIFSPGVRFPESCQDEVRDQKQLLKDAA AFLLSCQIPGLVKDCMEHAVLPVDGATLAEVMR QRGINMRYLGKVLELVLRSPARHQLDHVFKIGIG ELITRSAKHIFKTYLQGVELSGLSAAISHFLNCFLS SYPNVAHLPADELVSKKRNKRKNRPPGAADN TAWAVMTPQELWKNICQEAKNYFDDFDECETV DQAVETYGLQKITLLREISLKTGIQVLLKEYSFDS RHKPAFTEEDVLNIFPVVKHVNPKASDAFHFFQS GQAKVQQGFLKEGCELINEALNLFNNVYGAMH VETCACLRLLARLHYIMGDYAEALSNQQKAVL MSERVMGTEHPNTIQEYMHLALYCFASSQLSTA LSLLYRARYLMLLVFGEDHPEMALLDNNIGLVL HGVMEYDLSLRFLENALAVSTKYHGPKALKVAL SHHLVARVYESKAEFRSALQHEKEGYTIYKTQL GEDHEKTKESSEYLKCLTQQAVALQRTMNEIJYR NGSSANIPPLKFTAPSMASVLEQLNVINGILFIPLS QKDLENLKAEVARRHQLQEASRNRDRAEEPMA TEPAPAGAPGDLGSQPPAAKDPSPSVQG
3154	A	416	4082	KFKLIKIMLLTLIILLPVVSKFSFVSLSAPQHWSCP EGTLAGNGNSTCVGPAPFLIFSHGNSIFRIDTEGT NYEQLVVDAGVSVIMDFHYNEKRIYWVDLERQ LLQRVFLNGSRQERVCNIEKNVSGMAINWINEEV IWSNQQEGIITVTDMKGNNSHILLSALKYPANVA VDPVERFIFWSSEVAGSLYRADLDGVGVKALLE TSEKITAVSLDVLDKRLFWIQYNREGSNSLICSCD YDGGSVHISKHPTQHNLFAMSLFGDRIFYSTWK

II M ANIA I NG A

SEQ ID NO:	Method	Predicted beginning nucleotide location corresponding to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion
				MKTIWIANKHTGKDMVRINLHSSFVPLGELKVV HPLAQPKAEDDTWEPEQKLCKLRKGNCSSTVCG QDLQSHLCMCAEGYALSRDRKYCEGNDWKYCE DVNECAFWNHGCTLGCKNTPGSYYCTCPVGFVL LPDGKRCHQLVSCPRNVSECSHDCVLTSEGPLCF CPEGSVLERDGKTCSGCSSPDNGGCSQLCVPLSP VSWECDCFPGYDLQLDEKSCAASGPQPFLLFANS QDIRHMHFDGTDYGTLLSQQMGMVYALDHDPV ENKIYFAHTALKWIERANMDGSQRERLIEEGVD VPEGLAVDWIGRRFYWTDRGKSLIGRSDLNGKR SKIITIENISQPRGIAVHPMAKRLFWTDTGINPRIE SSSLQGLGRLVIASSDLIWPSGITIDFLTDKLYWC DAKQSVIEMANLDGSKRRRLTQNDVGHPFAVA VFEDYVWFSDWAMPSVIRVNKRTGKDRVRLQG SMLKPSSLVVVHPLAKPGADPCLYQNGGCEHIC KKRLGTAWCSCREGFMKASDGKTCLALDGHQL LAGGEVDLKNQVTPLDILSKTRVSEDNITESQHM LVAEIMVSDQDDCAPVGCSMYARCISEGEDATC QCLKGFAGDGKLCSDIDECEMGVPVCPPASSKCI NTEGGYVCRCSEGYQGDGIHCLDIDECQLGVHS CGENASCTNTEGGYTCMCAGRLSEPGLICPDSTP PPHLREDDHHYSVRNSDSECPLSHDGYCLHDGV CMYIEALDKYACNCVVGYIGERCQYRDLKWWE LRHAGHGQQQKVIVVAVCVVVLVMLLLLSLWG AHYYRTQKLLSKNPKNPYEESSRDVRSRRPADT EDGMSSCPQPWFVVIKEHQDLKNGGQPVAGED GQAADGSMQPTSWRQEPQLCGMGTEQGCWIPV SSDKGSCPQVMERSFHMPSYGTQTLEGGVEKPH
3155	A	533	212	SLLSANPLWQQRALDPPHQMELTQ GTSGWYWERLAERRGRLWSREEAMATMENKVI CALVLVSMLALGTLAEAQTETCTVAPRERQNCG FPGVTPSQCANKGCCFDDTVRGVPWCFYPNTID VPPEEECEF
3156	A	2	1585	PRVRAADVAAGAQAVVSAGMAKSNGENGPRAP AAGESLSGTRESLAQGPDAATTDELSSLGSDSEA NGFAERRIDKFGFIVGSQGAEGALEEVPLEVLRQ RESKWLDMLNNWDKWMAKKHKKIRLRCQKGI PPSLRGRAWQYLSGGKVKLQQNPGKFDELDMSP GDPKWLDVIERDLHRQFFFHEMFVSRGGHGQQD LFRVLKAYTLYRPEEGYCQAQAPIAAVLLMHMP AEQAFWCLVQICEKYLPGYYSEKLEAIQLDGEIL FSLLQKVSPVAHKHLSRQKIDPLLYMTEWFMCA FSRTLPWSSVLRVWDMFFCEGVKIIFRVGLVLLK HALGSPEKVKACQGQYETIERLRSLSPKIMQEAF LVQEVVELPVTERQIEREHLLQLRRWQETRGELQ CRSPPRLHGAKAILDAEPGPRPALQPSPSIRLPLD APLPGSKAKPKPPKQAQKEQRKQMKGRGQLEKP PAPNQAMVVAAAGDACPPQHVPPKDSAPKDSAP QDLAPQVSAHHRSQESLTSQESEDTYL
3157	A	3	601	SSAMGSRSSHAAVIPDGDSIRRETGFSQASLLRLH HRFRALDRNKKGYLSRMDLQQIGALAVNPLGDR IIESFFPDGSQRVDFPGFVRVLAHFRPVEDEDTET QDPKKPEPLNSRRNKLHYAFQLYDLDRDGKISR HEMLQVLRLMVGVQVTEEQLENIADRTVQEAD EDGDGAVSFVEFTKSLEKMDVEHKMSIRILK

SEQ ID NO:	Method	Predicted beginning nucleotide location corresponding to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon,/=possible nucleotide deletion, \=possible nucleotide insertion
3158	A	2	409	ISSCPHTAYEGSMSTLSNFTQTLEDVFRRIFITYM DNWRQNTTAEQEALQAKVDAENFYYVILYLMV MIGMFSFIIVAILVSTVKSKRREHSNDPYHQYIVE DWQEKYKSQILNLEESKATIHENIGAAGFKMSP
3159	A	3	416	PWGAAELDMGRRDAQLLAALLVLGLCALAGSE KPSPCQCSRLSPHNRTNCGFPGITSDQCFDNGCCF DSSVTGVPWCFHPLPKQESDQCVMEVSDRRNCG YPGISPEECASRKCCFSNFIFEVPWCFFPKSVEDC HY
3160	Α	179	409	KPKTKILKMVYYPELFVWVSQEPFPNKDMEGRL PKGRLPVPKEVNRKKNDETNAASLTPLGSSELRS PRISYLHFF
3161	A .	683	1186	LSSTGGLHAAACAAAMSLVIPEKFQHILRVLNTN IDGRRKIAFAITAIKGVGRRYAHVVLRKADIDLT KRAGELTEDEVERVITIMQNPRQYKIPDWFLNRQ KDVKDGKYSQVLANGLDNKLREDLERLKKIRA HRGLRHFWGLRVRGQHTKTTGRRGRTVGVSKK K
3162	A	1	1938	GMPRSRGGRAAPGPPPPPPPPPQQAPRWSRWRVP GRLLLLLPALCCLPGAARAAAAAAGAGNRAA VAVAVARADEAEAPFAGQNWLKSYGYLLPYDS RASALHSAKALQSAVSTMQQFYGIPVTGVLDQT TIEWMKKPRCGVPDHPHLSRRRRNKRYALTGQK WRQKHITYSIHNYTPKVGELDTRKAIRQAFDVW QKVTPLTFEEVPYHEIKSDRKEADIMIFFASGFHG DSSPFDGEGGFLAHAYFPGPGIGGDTHFDSDEPW TLGNANHDGNDLFLVAVHELGHALGLEHSSDPS AIMAPFYQYMETHNFKLPQDDLQGIQKIYGPPAE PLEPTRPLPTLPVRRIHSPSERKHERQPRPPRPPLG DRPSTPGTKPNICDGNFNTVALFRGEMFVFKDR WFWRLRNNRVQEGYPMQIEQFWKGLPARIDAA YERADGRFVFFKGDKYWVFKEVTVEPGYPHSLG ELGSCLPREGIDTALRWEPVGKTYFFKGERYWR YSEERRATDPGYPKPITVWKGIPQAPQGAFISKE GYYTYFYKGRDYWKFDNQKLSVEPGYPRNILRD WMGCNQKEVERRKERRLPQDDVDIMVTINDVP GSVNAVAVVIPCILSLCILVLVYTIFQFKNKTGPQ PVTYYKRPVQEWV
3163	A	1235	2223	SRLSLQFYVSFRRTGLFTCKLIVEIFFRNYMNDSL RTNVFVRFQPETIACACIYLAARALQIPLPTRPHW FLLFGTTEEEIQEICIETLRLYTRKKPNYELLEKEV EKRKVALQEAKLKAKGLNPDGTPALSTLGGFSP ASKPSSPREVKAEEKSPISINVKTVKKEPEDRQQA SKSPYNGVRKDSKRSRNSRSASRSRTRSRSRS HTPRRHYNNRRSRSGTYSSRSRSRSRSHSESPRR HHNHGSPHLKAKHTRDDLKSSNRHGHKRKKSRS RSQSKSRDHSDAAKKHRHERGHHRDRRERSRSF ERSHKSKHHGGSRSGHGRHRR
3164	A	3	3274	DCRLQAAMPTNFTVVPVEAHADGGGDETAERT EAPGTPEGPEPERPSPGDGNPRENSPFLNNVEVE QESFFEGKNMALFEEEMDSNPMVSSLLNKLANY TNLSQGVVEHEEDEESRRREAKAPRMGTFIGVY LPCLQNILGVILFLRLTWIVGVAGVLESFLIVAMC CTCTMLTAISMSAIATNGVVPAGGSYYMISRSLG PEFGGAVGLCFYLGTTFAGAMYILGTIEIFLTYISP

SEQ ID NO:	Méthod	Predicted beginning nucleotide location corresponding to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \possible nucleotide insertion
		·		GAAIFQAEAAGGEAAAMLHNMRVYGTCTLVLM ALVVFVGVKYVNKLALVFLACVVLSILAIYAGVI KSAFDPPDIPVCLLGNRTLSRRSFDACVKAYGIH NNSATSALWGLFCNGSQPSAACDEYFIQNNVTEI QGIPGAASGVFLENLWSTYAHAGAFVEKKGVPS VPVAEESRASTLPYVLTDIAASFTLLVGIYFPSVT GIMAGSNRSGDLKDAQKSIPTGTILAIVTTSFIYLS CIVLFGACIEGVVLRDKFGEALQGNLVIGMLAW PSPWVIVIGSFFSTCGAGLQTLTGAPRLLQAIARD GIVPFLQVFGHGKANGEPTWALLLTVLICETGILI ASLDSVAPILSMFFLMCYLFVNLACAVQTLLRTP NWRPRFKFYHWTLSFLGMSLCLALMFICSWYYA LSAMLIAGCIYKYIEYRGAEKEWGDGIRGLSLNA ARYALLRVEHGPPHTKNWRPQVLVMLNLDAEQ AMKHPRLLSFTSQLKAGKGLTIVGSVLEGTYLD KHMEAQRAEENIRSLMSTEKTKGFCQLVVSSSLR DGMSHLIQSAGLGGLKHNTVLMAWPASWKQED NPFSWKNFVDTVRDTTAAHQALLVAKNVDSFPQ NQERFGGGHIDVWWIVHDGGMLMLLPFLLRQH KVWRKCRMRIFTVAQVDDNSIQMKKDLQMFLY HLRISAEVEVVEMVENDISAFTYERTLMMEQRS QMLKQMQLSKNEQEREAQLIHDRNTASHTAAA ARTQAPPTPDKVQMTWTREKLIAEKYRSRDTSL SGFKDLFSMKPDQSNVRRMHTAVKLNGVVLNK SQDAQLVLLNMPGPPKNRQGDENYMEFLEVLTE GLNRVLLVRGGGREVITIYS
3165	A	3	2681	GRGARGGSGAGALRGCRGYLQKLSGKGPSRGY RSRWFVFDARRCYLYYFKSPQDALPLGHLDIAD ACFSYQGPDEAAEPGTEPPAHFQVHSAGAVTVL KAPNRQLMTYWLQELQQKRWEYCNSLDMVKW DSRTSPTPGDFPKGLVARDNTDLIYPHPNASAEK ARNVLAVETVPGELVGEQAANQPAPGHPNSINF YSLKQWGNELKNSMSSFRPGRGHNDSRRTVFYT NEEWELLDPTPKDLEESIVQEEKKKLTPEGNKGV TGSGFPFDFGRNPYKGKRPLKDIIGSYKNRHSSG DPSSEGTSGSGSVSIRKPASEMQLQVQSQQEELE QLKKDLSSQKELVRLLQQTVRSSQYDKYFTSSRL CEGVPKDTLELLHQKDDQILGLTSQLERFSLEKE SLQQEVRTLKSKVGELNEQLGMLMETIQAKDEV IIKLSEGEGNGPPPTVAPSSPSVVPVARDQLELDR LKDNLQGYKTQNKFLNKEILELSALRRNPERRER DLMARNSSLEAKLCQIESKYLILLQEMKTPVCSE DQGPTREVIAQLLEDALQVESQEQPEQAFVKPHL VSEYDIYGFRTVPEDDEEEKLVAKVRALDLKTL YLTENQEVSTGVKWENYFASTVNREMMCSPEL KNLIRAGIPHEHRSKVWK WCVDRHTRKFKDNTE PGHFQTLLQKALEKQNPASKQIELDLLRTLPNNK HYSCPTSEGIQKLRNVLLAFSWRNPDIGYCQGLN RLVAVALLYLEQEDAFWCLVTIVEVFMPRDYYT KTLLGSQVDQRVFRDLMSEKLPRLHGHFEQYKV DYTLITFNWFLVVFVDSVVSDILFKIWDSFLYEGP KVIFRFALALFKYKEEEILKLQDSMSIFKYLRYFT RTILDARSGTDAPTTWRKSGWS
3166	A	10	4070	FPGPTISSNSQLYRASALFETIRHEAQLSTDYKLS LFDLQTSSYQALQRVLVSLGHHDEALAVAERGR

SEQ ID NO:	Method	Predicted beginning nucleotide location corresponding to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion
				TRAFADLLVERQTGQQDSDPYSPVTIDQILEMVN GQRGLVLYYSLAAGYLYSWLLAPGAGIVKFHEH YLGENTVENSSDFQASSSVTLPTATGSALEQHIAS VREALGVESHYSRACASSETESEAGDIMDQGFEE MNNKLNSVTDPTGFLRMVRRNNLFNRSCQSMTS LFSNTVSPTQDGTSSLPRRQSSFAKPPLRALYDLL IAPMEGGLMHSSGPVGRHRQLILVLEGELYLIPF ALLKGSSSNEYLYERFGLLAVPSIRSLSVQSKSHL RKNPPTYSSSTSMAAVIGNPKLPSAVMDRWLWG PMPSAEEEAYMVSELLGCQPLVGSVATKERVMS ALTQAECVHFATHISWKLSALVLTPSMDGNPASS KSSFGHPYTIPESLRVQDDASDGESISDCPPLQEL LLTAADVLDLQLPVKLVVLGSSQESNSKVAADG VIALTRAFLAAGAQCVLVSLWPVPVAAFKMFIH AFYSSLLNGLKASAALGEAMKVVQSSKAFSHPS NWAGFMLIGSDVKLNSPSSLIGQALTEILQHPER ARDALRVLLHLVEKSLQRIQNGQRNAMYTSQQS VENKVGGIPGWQALLTAVGFRLDPPTSGLPAAV FFPTSDPGDRLQQCSSTLQSLLGLPNPALQALCK LITASETGEQLISRAVKNMVGMLHQVLVQLQAG EKEQDLASAPIQVSISVQLWRLPGCHEFLAALGF VLCEVGQEEVILKTGKQANRRTVHFALQSLLSLF DSTELPKRLSLDSSSSLESLASAQSVSNALPLGYQ QPPFSPTGADSIASDAISVYSLSSIASSMSFVSKPE GGSEGGGPGGRQDHDRSKNAYLQRSTLPRSQLP PQTRPAGNKDEEYEGFSIISNEPLATYQENRNTC FSPDHKQPQPGTAGGMRVSVSSKGSISTPNSPVK MTLIPSPNSPFQKVGKLASSDTGESDQSSTETDST VKSQEESNPKLDPQELAQKILEETQSHLIAVERLQ RSGGQVSKSNNPEDGVQAPSSTAVFRASETSAFS RPVLSHQKSQPSPVTVKPKPPARSSSLPKVSSGYS SPTTSEMSIKDSPSQHSGRPSPGCDSQTSQLDQPL FKLKYPSSPYSAHISKSPRNMSPSSGHQSPAGSAP SPALSYSSAGSARSSPADAPDIDKLKMAAIDEKV QAVHNLKMFWQSTPQHSTGPMKIFRGAPGTMTS KRDVLSLLNLSPRPNKKEEGVDKLELKELSLQQH DGAPPKAPPNGHWRTETTSLGSLPLPAGPPATAP ARPLRLPSGNGYKFLSPGRFFPSSKC
3167	A	1	762	AARRQKGKEENMMMDLFETGSYFFYLDGENV TLQPLEVAEGSPLYPGSDGTLSPCQDQMPPEAGS DSSGEEHVLAPPGLQPPHCPGQCLIWACKTCKRK SAPTDRRKAATLRERRRLKKINEAFEALKRRTVA NPNQRLPKVEILRSAISYIERLQDLLHRLDQQEK MQELGVDPFSYRPKQENLEGADFLRTCSSQWPS VSDHSRGLVITAKEGGASIDSSASSSLRCLSSIVDS ISSEERKLPCVEEVVEK
3168	A	701	246	TSRRVTMKFNPFVTSDRSKNRKRHFNAPSHVRR KIMSSPLSKELRQKYNVRSMPIRKDDEVQVVRG HYKGQQIGKVVQVYRKKYVIYIERVQREKANGT TVHVGIHPSKVVITRLKLDKDRKKILERKAKSRQ VGKEKGKYKEELIEKMQE
3169	A	156	3168	GPGGAISLSVEAKAGADLLVKGKQARMDIYDTQ TLGVVVFGGFMVVSAIGIFLVSTFSMKETSYEEA LANQRKEMAKTHHQKVEKKKKEKTVEKKGKT KKKEEKPNGKIPDHDPAPNVTVLLREPVRAPAV

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- 			Decal-4-3	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid,
SEQ ID NO:	Method	Predicted beginning	Predicted end nucleotide	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine,
NO:		nucleotide	location	I=Isoleucine, K=Lysine, L=Leucine, M=Methionine,
		location	corresponding	N=Asparagine, P=Proline, O=Glutamine, R=Arginine, S=Serine,
		corresponding	to last amino	T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine,
	1	to first amino	acid residue of	X=Unknown, *=Stop codon, /=possible nucleotide deletion,
		acid residue of	peptide	\=possible nucleotide insertion
		peptide sequence	sequence	
		sequence		AVAPTPVQPPIIVAPVATVPAMPQEKLASSPKDK
				KKKEKKVAKVEPAVSSVVNSIQVLTSKAAILETA
ļ	[PKEGRNTDVAQSPEAPKQEAPAKKKSGSKKKGP
				PDADGPLYLPYKTLVSTVGSMVFNEGEAQRLIEI
				LSEKAGIIQDTWHKATQKGDPVAILKRQLEEKEK
		i .		LLATEQEDAAVAKSKLRELNKEMAAEKAKAAA
	l	į		GEAKVKKQLVAREQEITAVQARMQASYREHVK
				EVQQLQGKIRTLQEQLENGPNTQLARLQQENSIL
	1			
				RDALNQATSQVESKQNAELAKLRQELSKVSKEL
		1		VEKSEAVRQDEQQRKALEAKAAAFEKQVLQLQ
			İ	ASHRESEEALQKRLDEVSRELCHTQSSHASLRAD
]			AEKAQEQQQMAELHSKLQSSEAEVRSKCEELS
				GLHGQLQEARAENSQLTERIRSIEALLEAGQARD
				AQDVQASQAEADQQQTRLKELESQVSGLEKEAI
				ELREAVEQQKVKNNDLREKNWKAMEALATAEQ
	1			ACKEKLHSLTQAKEESEKQLCLIEAQTMEALLAL
1	ľ			LPELSVLAQQNYTEWLQDLKEKGPTLLKHPPAP
İ				AEPSSDLASKLREAEETQSTLQAECDQYRSILAET
				EGMLRDLQKSVEEEEQVWRAKVGAAEEELQKS
				RVTVKHLEEIVEKLKGELESSDQVREHTSHLEAE
ļ	•			LEKHMAAASAECQNYAKEVAGLRQLLLESQSQL
				DAAKSEAQKQSDELALVRQQLSEMKSHVEDGDI
İ				AGAPASSPEAPPAEQDPVQLKTQLEWTEAILEDE
		1		QTQRQKLTAEFEEAQTSACRLQEELEKLRTAGPL
			1	ESSETEEASQLKERLEKEKKLTSDLGRAATRLQE
		ļ		LLKTTQEQLAREKDTVKKLQEQLEKAEDGSSSK
				EGTSV
3170	Α .	6730	4027	THASEKYSYGHLPTHSITAHPMVTIRISDRQRLIQ
3170	1.	0,30		PYIHNYSWLLFAALALYSAHLASAEDVDGEKLD
		}		POTRSSATTLRSQCMQLVGDCLMKAHQGKGLK
				ALALLGVLPDGDSSLEDHALPVTVPTGASEEQLE
				KKAVQGAELSEAGNGKRAVHEEIRPVDFKQRNK
ļ		ļ	Ì	ADKGVSLSKDPSCQTQISDSPADASPPTGLPDAE
			{	DSEVSSOKPIEEKAVTPSPEQVFAECSQKRILGLL
<u> </u>				AAMLPPLKSGPTVPLIDLEHVLPLMFQVVISNAG
				HLNETYHLTLGLLGQLIIRLLPAEVDAAVIKVLSA
		İ		KHNLFAAGDSSIVPDGWKTTHLLFSLGAVCLDS
		1		RVGLDWACSMAEILRSLNSAPLWRDVIATFTDH
		1		CIKQLPFQLKHTNIFTLLVLVGFPQVLCVGTRCV
		1]	YMDNANEPHNVIILKHFTEKNRAVIVDVKTRKR
		1		KTVKDYQLVQKGGGQECGDSRAQLSQYSQHFA
		1		FIASHLLQSSMDSHCPEAVEATWVLSLALKGLY
			· .	KTLKAHGFEEIRATFLQTDLLKLLVKKCSKGTGF
}		1		SKTWLLRDLEILSIMLYSSKKEINALAEHGDLEL
		1		DERGDREEEVERPVSSPGDPEQKKLDPLEGLDEP
1				TRICFLMAHDALNAPLHILRAIYELQMKKTDYFF
}		1		LEVQKRFDGDELTTDERIRSLAQRWQPSKSLRLE
1			1	EQSAKAVDTDMIILPCLSRPARCDQATAESNPVT
1		1	1	QKLISSTESELQQSYAKQRRSKSAALLHKELNCK
		1		SKRAVRDYLFRVNEATAVLYARHVLASLLAEWP
1				SHVPVSEDILELSGPAHMTYILDMFMQLEEKHE
l ·		1	[WEKVVMQTELVLTHQVLPLPHRLPPVSASWSEA
l				TCVAVQLPDRCECSKGRVTVSSPKDWASEELRG
1				
	i		1	
3171	A	557	89	PERDFQLNQKALSPSSQFPSAEILRHIR GTRAGPVKDREAFQRLNFLYQAAHCVLAQDPEN

		T-8-11-	[N 31.4 N	LA -t
SEQ ID NO:	Method	Predicted	Predicted end nucleotide	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine,
NO:		beginning nucleotide	location	I=Isoleucine, K=Lysine, L=Leucine, M=Methionine,
		location	corresponding	N=Asparagine, P=Proline, O=Glutamine, R=Arginine, S=Serine,
		corresponding	to last amino	T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine,
		to first amino	acid residue of	X=Unknown, *=Stop codon, /=possible nucleotide deletion,
		acid residue of peptide sequence	peptide sequence	
	<u> </u>	sequence		QALARFYCYTERTIAKRLVLRRDPSVKRTLCRGC
				SSLLVPGLTCTQRQRRCRGQRWTVQTCLTCQRS
				QRFLNDPGHLLWGDRPEAQLGSQADSKPLQPLP
				NTAHSISDRLPEEKMQTQGSSNQ
3172	A	2	496	FRRAGAGRGRRRGEVTSPLSPEPLAFQSLATSRR
				PEPQTTQTVRSSALPAPPASPMSQYAPSPDFKRA
		İ		LDSSPEANTEDDKTEEDVPMPKNYLWLTIVSCFC
				PAYPINIVALVFSIMSLNSYNDGDYEGARRLGRN
	<u></u>		4040	AKWVAIASIIIGLLIIGISCAVHFTRNA
3173	Α	2	4048	FRSGGCRRAWTSRWPQRRRSPESCEAPLSAPL WGPORGLPGREPLRSRSASAIALRTIGHILALLLR
				LLHLGLGSGGCREDVPPSGRGKKEEKMKKHRRA
				LALVSCLFLCSLVWLPSWRVCCKESSSASASSYY
		1		SQDDNCALENEDVQFQKKDEREGPINAESLGKS
1				GSNLPISPKEHKLKDDSIVDVQNTESKKLSPPVVE
		ĺ	[TLPTVDLHEESSNAVVDSETVENISSSSTSEITPIS
				KLDEIEKSGTIPIAKPSETEQSETDCDVGEALDAS
				APIEQPSFVSPPDSLVGQHIENVSSSHGKGKITKSE
				FESKVSASEQGGGDPKSALNASDNLKNESSDYT
				KPGDIDPTSVASPKDPEDIPTFDEWKKKVMEVEK
				EKSQSMHASSNGGSHATKKVQKNRNNYASVEC
1		1		GAKILAANPEAKSTSAILIENMDLYMLNPCSTKI
		1		WFVIELCEPIQVKQLDIANYELFSSTPKDFLVSISD
				RYPTNKWIKLGTFHGRDERNVQSFPLDEQMYAK
				YVKMFIKYIKVELLSHFGSEHFCPLSLIRVFGTSM
				VEEYEEIADSQYHSERQELFDEDYDYPLDYNTGE
	· ·		+	DKSSKNLLGSATNAILNMVNIAANILGAKTEDLT
				EGNKSISENATATAAPKMPESTPVSTPVPSPEYVT
		•		TEVHTHDMEPSTPDTPKESPIVQLVQEEEEEASPS
				TVTLLGSGEQEDESSPWFESETQIFCSELTTICCIS
	İ			SFSEYIYKWCSVRVALYRQRSRTALSKGKDYLV
				LAQPPLLLPAESVDVSVLQPLSGELENTNIEREAE TVVLGDLSSSMHQDDLVNHTVDAVELEPSHSQT
		ĺ		LSQSLLLDITPEINPLPKIEVSESVEYEAGHIPSPVI
,				PQESSVEIDNETEQKSESFSSIEKPSITYETNKVNE
				LMDNIIKEDVNSMQIFTKLSETIVPPINTATVPDN
				EDGEAKMNIADTAKQTLISVVDSSSLPEVKEEEQ
				SPEDALLRGLQRTATDFYAELQNSTDLGYANGN
	ĺ			LVHGSNQKESVFMRLNNRIKALEVNMSLSGRYL
		1		EELSQRYRKQMEEMQKAFNKTIVKLQNTSRIAE
				EQDQRQTEAIQLLQAQLTNMTQLVSNLSATVAE
				LKREVSDRQSYLVISLVLCVVLGLMLCMQRCRN
				TSQFDGDYISKLPKSNQYPSPKRCFSSYDDMNLK
	1			RRTSFPLMRSKSLQLTGKEVDPNDLYIVEPLKFSP
1				EKKKKRCKYKIEKIETIKPEEPLHPIANGDIKGRK
				PFTNQRDFSNMGEVYHSSYKGPPSEGSSETSSQS
				EESYFCGISACTSLCNGQSQKTKTEKRALKRRRS
1				KVQDQGKLIKTLIQTKSGSLPSLHDIIKGNKEITV
	<u> </u>		1	GTFGVTAVSGHI
3174	A	485	4668	RKCSKEKASKTPSQKIPTTPCCVLQAGPEPRSLAE
				RMGADGETVVLKNMLIGVNLILLGSMIKPSECQL
1		i .	1	EVTTERVQRQSVEEEGGIANYNTSSKEQPVVFNH
				VYNINVPLDNLCSSGLEASAEQEVSAEDETLAEY MGQTSDHESQVTFTHRINFPKKACPCASSAQVLQ
				ELLSRIEMLEREVSVLRDQCNANCCQESAATGQL
L	L		<u> </u>	PETPICIFICE A 2 A PICT A CHARACTER WATER

SEQ ID NO:	Method	Predicted beginning	Predicted end	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine,
NO:		nucleotide	location	I=Isoleucine, K=Lysine, L=Leucine, M=Methionine,
		location	corresponding	N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine,
		corresponding to first amino	to last amino acid residue of	T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion,
		acid residue of	peptide	
		peptide sequence	sequence	
				DYIPHCSGHGNFSFESCGCICNEGWFGKNCSEPY CPLGCSSRGVCVDGQCICDSEYSGDDCSELRCPT
1				DCSSRGLCVDGECVCEEPYTGEDCRELRCPGDCS
				GKGRCANGTCLCEEGYVGEDCGQRQCLNACSG
				RGQCEEGLCVCEEGYQGPDCSAVAPPEDLRVAG
	1			ISDRSIELEWDGPMAVTEYVISYQPTALGGLQLQ
1		1	}	QRVPGDWSGVTITELEPGLTYNISVYAVISNILSL
		:	İ	PITAKVATHLSTPQGLQFKTITETTVEVQWEPFSF
		1		SFDGWEISFIPKNNEGGVIAQVPSDVTSFNQTGLK
				PGEEYIVNVVALKEQARSPPTSASVSTVIDGPTQI
				LVRDVSDTVAFVEWIPPRAKVDFILLKYGLVGGE
				GGRTTFRLQPPLSQYSVQALRPGSRYEVSVSAVR GTNESDSATTQFTTEIDAPKNLRVGSRTATSLDL
				EWDNSEAEVQEYKVVYITLAGEQYHEVLVPRGI
	,			GPTTRATLTDLVPGTEYGVGISAVMNSQQSVPAT
				MNARTELDSPRDLMVTASSETSISLIWTKASGPID
}		1		HYRITFTPSSGIASEVTVPKDRTSYTLTDLEPGAE
				YIISVTAERGRQQSLESTVDAFTGFRPISHLHFSH
ľ		1	ĺ	VTSSSVNITWSDPSPPADRLILNYSPRDEEEEMME
	:			VSLDATKRHAVLMGLQPATEYIVNLVAVHGTVT
				SEPIVGSITTGIDPPKDITISNVTKDSVMVSWSPPV
				ASFDYYRVSYRPTQVGRLDSSVVPNTVTEFTITR LNPATEYEISLNSVRGREESERICTLVHTAMDNP
		·Ω		VDLIATNITPTEALLQWKAPVGEVENYVIVLTHF
				AVAGETILVDGVSEEFRLVDLLPSTHYTATMYAT
				NGPLTSGTISTNFSTLLDPPANLTASEVTRQSALIS
				WQPPRAEIENYVLTYKSTDGSRKELIVDAEDTWI
				RLEGLLENTDYTVLLQAAQDTTWSSITSTAFTTG
				GRVFPHPQDCAQHLMNGDTLSGVYPIFLNGELS
		}		QKLQVYCDMTTDGGGWIVFQRRQNGQTDFFRK WADYRVGFGNVEDEFWLGLDNIHRITSQGRYEL
	'			RVDMRDGQEAAFASYDRFSVEDSRNLYKLRIGS
				YNGTAGDSLSYHQGRPFSTEDRDNDVAVTNCA
		ł		MSYKGAWWYKNCHRTNLNGKYGESRHSQGIN
				WYHWKGHEFSIPFVEMKMRPYNHRLMAGRKRQ
				SLQF
3175	A	2	623	RLQLPACPALSAAHPLALPSFSSQCHRAEARAAA
				AATAEGTMASGVTVNDEVIKVFNDMKVRKSST
				QEEIKKRKKAVLFCLSDDKRQIIVEEAKQILVGDI GDTVEDPYTSFVKLLPLNDCRYALYDATYETKE
				SKKEDLVFIFWAPESAPLKSKMIYASSKDAIKKK
				FTGIKHEWQVNGLDDIKDRSTLGEKLGGNVVVS
				LEGKPL
3176	Α	99	1567	PRGCWSSCLDAMFRLNSLSALAELAVGSRWYH
	1			GGSQPIQIRRRLMMVAFLGASAVTASTGLLWKR
				AHAESPPCVDNLKSDIGDKGKNKDEGDVCNHEK
				KTADLAPHPEEKKKKRSGFRDRKVMEYENRIRA
		ļ		YSTPDKIFRYFATLKVISEPGEAEVFMTPEDFVRS
				ITPNEKQPEHLGLDQYIIKRFDGKTEKISQEREKF ADEGSIFYTLGECGLISFSDYIFLTTVLSTPQRNFE
		}		IAFKMFDLNGDGEVDMEEFEQVQSIIRSQTSMG
			ļ	MRHRDRPTTGNTLKSGLCSALTTYFFGADLKGK
				LTIKNFLEFQRKLQHDVLKLEFERHDPVDGRITE
				RQFGGMLLAYSGVQSKKLTAMQRQLKKHFKEG
L	<u></u>	<u> </u>		KGLTFQEVENFFTFLKNINDVDTALSFYHMAGAS

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SEQ ID NO:	Method	Predicted beginning nucleotide location corresponding to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methlonine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion
				LDKVTMQQVARTVAKVELSDHVCDVVFALFDC DGNGELSNKEFVSIMKQRLMRGLEKPKDMGFTR LMQAMWKCAQETAWDFALPKQ
3177	A	182	648	LGVVGSGAAVGGRQAARGAALGRRPMAAVLG ALGATRRLLAALRGQSLGLAAMSSGTHRLTAEE RNQAILDLKAAGWSELSERDAIYKEFSFHNFNQA FGFMSRVALQAEKMNHHPEWFNVYNKVQITLTS HDCGELTKKDVKLAKFIEKAAASV
3178	A	8	612	ACGCRSFCGSTVMSLLLYYALPALGSYAMLSIFF LRRPHLLHTPRAPTFRIRLGAHRGGSGELLENTM EAMENSMAQRSDLLELDCQLTRDRVVVVSHDE NLCRQSGLNRDVGSLDFEDLPLYKEKLEVYFSPG HFAHGSDRRMVRLEDLFQRFPRTPMSVEIKGKN EELIREIAGLVRRYDRNEITIWASEKSSVMKKCK
3179	A	88	1496	QETSKMETLSFPRYNVAEIVIHIRNKILTGADGKN LTKNDLYPNPKPEVLHMIYMRALQIVYGIRLEHF YMMPVNSEVMYPHLMEGFLPFSNLVTHLDSFLPI CRVNDFETADILCPKAKRTSRFLSGIINFIHFREAC RETYMEFLWQYKSSADKMQQLNAAHQEALMK LERLDSVPVEEQEEFKQLSDGIQELQQSLNQDFH QKTIVLQEGNSQKKSNISEKTKRLNELKLSVVSL KEIQESLKTKIVDSPEKLKNYKEKMKDTVQKLK NARQEVVEKYEIYGDSVDCLPSCQLEVQLYQKK IQDLSDNREKLASILKESLNLEDQIESDESELKKL KTEENSFKRLMIVKKEKLATAQFKINKKHEDVK QYKRTVIEDCNKVQEKRGAVYERVTTINHEIQKI RLGIQQLKDAADREKLKSQEIFLNLKTALEKYHD GIEKAAEDSYAKIDEKTAELKRKMFKMST
	A		7086	GNMACWPQLRLLLWKNLTFRRRQTCQLLLEVA WPLFIFLILISVRLSYPPYEQHECHFPNKAMPSAG TLPWVQGIICNANNPCFRYPTPGEAPGVVGNFNK SIVARLFSDARRLLLYSQKDTSMKDMRKVLRTL QQIKKSSSNLKLQDFLVDNETFSGFLYHNLSLPK STVDKMLRADVILHKVFLQGYQLHLTSLCNGSK SEEMIQLGDQEVSELCGLPREKLAAAERVLRSN MDILKPILRTLNSTSPFPSKELAEATKTLLHSLGT LAQELFSMRSWSDMRQEVMFLTNVNSSSSSTQI YQAVSRIVCGHPEGGGLKIKSLNWYEDNNYKAL FGGNGTEEDAETFYDNSTTPYCNDLMKNLESSPL SRIIWKALKPLLVGKILYTPDTPATRQVMAEVNK TFQELAVFHDLEGMWEELSPKIWTFMENSQEMD LVRMLLDSRDNDHFWEQQLDGLDWTAQDIVAF LAKHPEDVQSSNGSVYTWREAFNETNQAIRTISR FMECVNLNKLEPIATEVWLINKSMELLDERKFW AGIVFTGITPGSIELPHHVKYKIRMGIDNVERTNK IKDGYWDPGPRADPFEDMRYVWGGFAYLQDVV EQAIIRVLTGTEKKTGVYMQQMPYPCYVDDIFLR VMSRSMPLFMTLAWIYSVAVIIKGIVYEKEARLK ETMRIMGLDNSILWFSWFISSLIPLLVSAGLLVVI LKLGNLLPYSDPSVVFVFLSVFAVVTILQCFLIST LFSRANLAAACGGIIYFTLYLPYVLCVAWQDYV GFTLKIFASLLSPVAFGFGCEYFALFEEQGIGVQW DNLFESPVEEDGFNLTTSVSMMLFDTFLYGVMT WYIEAVFPGQYGIPRPWYFPCTKSYWFGEESDEK SHPGSNQKRISEICMEEEPTHLKLGVSIQNLVKVY

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	r	l m	Dwadieted and	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid,
SEQ ID	Method	Predicted beginning	Predicted end nucleotide	E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine,
NO:		nucleotide	location	I=Isoleucine, K=Lysine, L=Leucine, M=Methionine,
	ŀ	location	corresponding	N=Asparagine, P=Proline, O=Glutamine, R=Arginine, S=Serine,
		corresponding	to last amino	T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine,
	•	to first amino	acid residue of	X=Unknown, *=Stop codon, /=possible nucleotide deletion,
		acid residue of peptide	peptide sequence	\=possible nucleotide insertion .
		sequence	sequence	
· <u> </u>	-	 		RDGMKVAVDGLALNFYEGQITSFLGHNGAGKTT
		i		TMSILTGLFPPTSGTAYILGKDIRSEMSTIRQNLG
				VCPQHNVLFDMLTVEEHIWFYARLKGLSEKHVK
				AEMEQMALDVGLPSSKLKSKTSQLSGGMQRKLS
				VALAFVGGSKVVILDEPTAGVDPYSRRGIWELLL
				KYRQGRTIILSTHHMDEADVLGDRIAIISHGKLCC
]	VGSSLFLKNQLGTGYYLTLVKKDVESSLSSCRNS
				SSTVSYLKKEDSVSQSSSDAGLGSDHESDTLTID
			†	VSAISNLIRKHVSEARLVEDIGHELTYVLPYEAA
			}	KEGAFVELFHEIDDRLSDLGISSYGISETTLEEIFL
			ŀ	KVAEESGVDAETSDGTLPARRNRRAFGDKQSCL
	1		1	RPFTEDDAADPNDSDIDPESRETDLLSGMDGKGS
			1	YQVKGWKLTQQQFVALLWKRLLIARRSRKGFF
			1	AQIVLPAVFVCIALVFSLIVPPFGKYPSLELQPWM
	1			YNEQYTFVSNDAPEDTGTLELLNALTKDPGFGT
				RCMEGNPIPDTPCQAGEEEWTTAPVPQTIMDLFQ
				NGNWTMQNPSPACQCSSDKIKKMLPVCPPGAGG
				LPPPQRKQNTADILQDLTGRNISDYLVKTYVQIIA
				KSLKNKIWVNEFRYGGFSLGVSNTQALPPSQEV
				NDATKOMKKHLKLAKDSSADRFLNSLGRFMTG
				LDTRNNVKVWFNNKGWHAISSFLNVINNAILRA
		j		NLOKGENPSHYGITAFNHPLNLTKQQLSEVAPM
				TTSVDVLVSICVIFAMSFVPASFVVFLIQERVSKA
				KHLQFISGVKPVIYWLSNFVWDMCNYVVPATLV
			,	IIIFICFQQKSYVSSTNLPVLALLLLLYGWSITPLM
				YPASFVFKIPSTAYVVLTSVNLFIGINGSVATFVL
				ELFTDNKLNNINDILKSVFLIFPHFCLGRGLIDMV
				KNQAMADALERFGENRFVSPLSWDLVGRNLFA
				MAVEGVVFFLITVLIQYRFFIRPRPVNAKLSPLND
				EDEDVRRERQRILDGGGQNDILEIKELTKIYRRK
	,			RKPAVDRICVGIPPGECFGLLGVNGAGKSSTFKM
				LTGDTTVTRGDAFLNRNSILSNIHEVHQNMGYCP
				QFDAITELLTGREHVEFFALLRGVPEKEVGKVGE
			•	WAIRKLGLVKYGEKYAGNYSGGNKRKLSTAMA
				LIGGPPVVFLDEPTTGMDPKARRFLWNCALSVV
				KEGRSVVLTSHSMEECEALCTRMAIMVNGRFRC
				LGSVQHLKNRFGDGYTIVVRIAGSNPDLKPVQDF
		i		FGLAFPGSVPKEKHRNMLQYQLPSSLSSLARIFSI
				LSQSKKRLHIEDYSVSQTTLDQVFVNFAKDQSDD
		İ		DHLKDLSLHKNQTVVDVAVLTSFLQDEKVKESY
				V
3181	A	215	1367	PPATSQAALPEALSKGRETPRPATHPARSQDVRP
2101	A		1307	LSCPFDFLRDNVEWSEEQAAAAERKVQENSIQR
,				VCQEKQVDYEINAHKYWNDFYKIHENGFFKDR
				HWLFTEFPELAPSQNQNHLKDWFLENKSEVPEC
				RNNEDGPGLIMEEQHKCSSKSLEHKTQTPPVEEN
		1	1	VTQKISDLEICADEFPGSSATYRILEVGCGVGNTV
			ļ	FPILOTNNDPGLFVYCCDFSSTAIELVQTNSEYDP
			(
				SRCFAFVHDLCDEEKSYPVPKGSLDIILIFVLSAI
			1	VPDKMQKAINRLSRLLKPGGMVLLRDYGRYDM
				AQLRFKKGQCLSGNFYVRGDGTRVYFFTQEELD
				TLFTTAGLEKVQNLVDRRLQVNRGKQLTMYRV
0100	ļ.,	1	1000	WIQCKYCKPLLSSTS
3182	A	3	1289	GSETQHLPRDPQHLPWDPQQHQDRRPELFHAF
			I	ARDSAPPPSMVLAAETTSQQERLQAIAEKRKRQ

CONO VD	T NG-02 - 3	130.131.4.3	Dundistad and	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid,
SEQ ID NO:	Method	Predicted beginning nucleotide location corresponding	Predicted end nucleotide location corresponding to last amino	E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine,
		to first amino acid residue of peptide sequence	acid residue of peptide sequence	X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion
				AEIENKRRQLEDERRQLQHLKSKALRERWLLEG
				TPSSASEGDEDLRRQMQDDEQKTRLLEDSVSRLE KGIEVLERGDSAPAAAKENAAAPSPVRAPAPSPA
				KEERKTEVVMNSQQTPVGTPKDKRVSNTPLRTV
				DGSPMMKAAMYSVEITVEKDKVTGETRVLSSTT
		į		LLPRQPLPLGIKVYEDETKVVHAVDGTAENGIHP
		-		LSSSEVDELIHKADEVTLSEAGSTAGAAETRGAV EGAARTTPSRREITGVQAQPGEATSGPPGIQPGQE
				PPVTMIFMGYQNVEDEAETKKVLGLQDTITAEL
				VVIEDAAEPKEPAPPNGSAAEPPTEAASREENQA
	<u> </u>		1001	GPEATTSDPQDLDMKKHRCKCCSIM
3183	A	333	1931	IAPTGGSHSEIQKQLGSGGDSSSQRRAERRTEPRS APRPRWGRSARSPGAHKLPGPPRRRDPGAWARL
		-		EAAAAHRHSRGSMGRRMRGAAATAGLWLLAL
				GSLLALWGGLLPPRTELPASRPPEDRLPRRPARS
				GGPAPAPRFPLPPPLAWDARGGSLKTFRALLTLA AGADGPPRQSRSEPRWHVSARQPRPEESAAVHG
				GVFWSRGLEEQVPPGFSEAQAAAWLEAARGAR
				MVALERGGCGRSSNRLARFADGTRACVRYGINP
				EQIQGEALSYYLARLLGLQRHVPPLALARVEAR GAQWAQVQEELRAAHWTEGSVVSLTRWLPNLT
				DVVVPAPWRSEDGRLRPLRDAGGELANLSQAEL
				VDLVQWTDLILFDYLTANFDRLVSNLFSLQWDP
				RVMQRATSNLHRGPGGALVFLDNEAGLVHGYR
				VAGMWDKYNEPLLQSVCVFRERTARRVLELHR . GQDAAARLLRLYRRHEPRFPELAALADPHAQLL
				QRRLDFLAKHILHCKAKYGRRSGDLVSPGGKER
				DLGLGYG
3184	A	1	1004 .	GSTHASADAWAQWFCTEALVMGAPVWYLVAA ALLVGFILFLTRSRGRAASAGQEPLHNEELAGAG
				RVAQPGPLEPEEPRAGGRPRRRDLGSRLQAQR
	-			RAQRVAWAEADENEEEAVILAQEEEGVEKPAET
			ļ	HLSGKIGAKKLRKLEEKQARKAQREAEEAEREE RKRLESQREAEWKKEEERLRLEEEQKEEEERKA
				REEQAQREHEEYLKLKEAFVVEEEGVGETMTEE
			1	QSQSFLTEFINYIKQSKVVLLEDLASQVGLRTQD
			1	TINRIQDLLAEGTITGVIDDRGKFIYITPEELAAVA NFIRQRGRVSIAELAQASNSLIAWGRESPAQAPA
3185	A	2981	7173	CLLAGKFSSTLYETGGCDMSLVNFEPAARRASNI
3103	1			CDTDSHVSSSTSVRFYPHDVLSLPQIRLNRLLTID
				TDLLEQQDIDLSPDLAATYGPTEEAAQKVKHYY
				RFWILPQLWIGINFDRLTLLALFDRNREILENVLA VILAILVAFLGSILLIQGFFRDIWVFQFCLVIASCQ
				YSLLKSVQPDSSSPRHGHNRIIAYSRPVYFCICCG
				LIWLLDYGSRNLTATKFKLYGITFTNPLVFISARD
				LVIVFTLCFPIVFFIGLLPQVNTFVMYLCEQLDIHI FGGNATTSLLAALYSFICSIVAVALLYGLCYGAL
				KDSWDGQHIPVLFSIFCGLLVAVSYHLSRQSSDP
!				SVLFSLVQSKIFPKTEEKNPEDPLSEVKDPLPEKL
				RNSVSERLQSDLVVCIVIGVLYFAIHVSTVFTVLQ
				PALKYVLYTLVGFVGFVTHYVLPQVRKQLPWH CFSHPLLKTLEYNQYEVRNAATMMWFEKLHVW
				LLFVEKNIIYPLIVLNELSSSAETIASPKKLNTELG
				ALMITVAGLKLLRSSFSSPTYQYVTVIFTVLFFKF
L		1	I	DYEAFSETMLLDLFFMSILFNKLWELLYKLQFVY

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SEQ ID NO:	Method	Predicted beginning nucleotide location corresponding to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \possible nucleotide insertion
				TYIAPWQITWGSAFHAFAQPFAVPHSAMLFIQAA VSAFFSTPLNPFLGSAIFITSYVRPVKFWERDYNT KRVDHSNTRLASQLDRNPGTYCQQREVEAITEG VEEDEGFCCCEPGHIPHMLSFNAAFSQRWLAWE VIVTKYILEGYSITDNSAASMLQVFDLRKVLTTY YVKGIIYYVTTSSKLEEWLANETMQEGLRLCAD RNYVDVDPTFNPNIDEDYDHRLAGISRESFCVIY LNWIEYCSSRRAKPVDVDKDSSLVTLCYGLCVL GRRALGTASHHMSSNLESFLYGLHALFKGDFRIS SIRDEWIFADMELLRKVVVPGIRMSIKLHQDHFT SPDEYDDPTVLYEAIVSHEKNLVIAHEGDPAWRS AVLANSPSLLALRHVMDDGTNEYKIIMLNRRYL SFRVIKVNKECVRGLWAGQQQELVFLRNRNPER GSIQNAKQALRNMINSSCDQPIGYPIFVSPLTTSY SDSHEQLKDILGGPISLGNIRNFIVSTWHRLRKGC GAGCNSGGNIEDSDTGGGTSCTGNNATTANNPH SNVTQGSIGNPGQGSGTGLHPPVTSYPPTLGTSHS SHSVQSGLVRQSPARASVASQSSYCYSSRHSSLR MSTTGFVPCRRSSTSQISLRNLPSSIQSRLSMVNQ MEPSGQSGLACVQHGLPSSSSSSQSIPACKHHTL VGFLATEGGQSSATDAQPGNTLSPANNSHSRKA EVIYRVQIVDPSQILEGINLSKRKELQWPDEGIRL KAGRNSWKDWSPQEGMEGHVIHRWVPCSRDPG TRSHIDKAVLLVQIDDKYVTVIETGVLELGAEV
3186	A	3	470	SLSAMRFLAATFLLLALSTAAQAEPVQFKDCGSV DGVIKEVNVSPCPTQPCQLSKGQSYSVNVTFTSN IQSKSSKAVVHGILMGVPVPFPIPEPDGCKSGINC PIQKDKTYSYLNKLPVKSEYPSIKLVVEWQLQDD KNQSLFCWEIPVQIVSHL
3187	A	3	470	SLSAMRFLAATFLLLALSTAAQAEPVQFKDCGSV DGVIKEVNVSPCPTQPCQLSKGQSYSVNVTFTSN IQSKSSKAVVHGILMGVPVPFPIPEPDGCKSGINC PIQKDKTYSYLNKLPVKSEYPSIKLVVEWQLQDD KNQSLFCWEIPVQIVSHL
3188	A		3483	PRVRTKLILLVNDKKRYERVGGGPKRLGRDVEM EEMIEQLQEKVHELEKQNDTLKNRLISAKQQLQT QGYRQTPYNNVQSRINTGRRKANENAGLQECPR KGIKFQDADVAETPHPMFTKYGNSLLEEARGEIR NLENVIQSQRGQIEELEHLAEILKTQLRRKENEIE LSLLQLREQQATDQRSNIRDNVEMIKLHKQLVE KSNALSAMEGKFIQLQEKQRTLKISHDALMANG DELNMQLKEQRLKCCSLEKQLHSMKFSERRIEEL QDRINDLEKERELLKENYDKLYDSAFSAAHEEQ WKLKEQQLKVQIAQLETALKSDLTDKTEILDRL KTERDQNEKLVQENRELQLQYLEQKQQLDELKK RIKLYNQENDINADELSEALLLIKAQKEQKNGDL SFLVKVDSEINKDLERSMRELQATHAETVQELEK TRNMLIMQHKINKDYQMEVEAVTRKMENLQQD YELKVEQYVHLLDIRAARIHKLEAQLKDIAYGTK QYKFKPEIMPDDSVDEFDETIHLERGENLFEIHIN KVTFSSEVLQASGDKEPVTFCTYAFYDFELQTTP VVRGLHPEYNFTSQYLVHVNDLFLQYIQKNTITL EVHQAYSTEYETIAACQLKFHEILEKSGRIFCTAS LIGTKGDIPNFGTVEYWFRLRVPMDQAIRLYRER AKALGYITSNFKGPEHMQSLSQQAPKTAQLSSTD

SEQ ID NO:	Method	Predicted beginning nucleotide location corresponding to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion
				STDGNLNELHITIRCCNHLQSRASHLQPHPYVVY KFFDFADHDTAIIPSSNDPQFDDHMYFPVPMNM DLDRYLKSESLSFYVFDDSDTQENIYIGKVNVPLI SLAHDRCISGIFELTDHQKHPAGTIHVILKWKFA YLPPSGSITTEDLGNFIRSEEPEVVQRLPPASSVST LVLAPRPKPRQRLTPVDKKVSFVDIMPHQSDVSQ EGSVDEVKENTEKMQQGKDDVSLLSEGQLAEQS LASSEDETEITEDLEPEVEEDMSASDSDDCIIPGPI SKNIKQPSEKIRIEIIALSLNDSQVTMDDTIQRLFV ECRFYSLPAEETPVSLPKPKSGQWVYYNYSNVIY VDKENNKAKRDILKAILQKQEMPNRSLRFTVVS DPPEDEQDLECEDIGVAHVDLADMFQEGRDLIE QNIDVFDARADGEGIGKLRVTVEALHALQSVYK QYRDDLEA
3189	A	476	1175	MKGSGWHLRSGMVGTLITTILPHWRRTAHVGTN ILTAVSYLKGLWMECVWHSTGIYQCQIYRSLLA LPQDLQAARALMGISCLLSGIACACAVIGMKCTR CAKGTPAKTTFAILGGTLFILAGLLCMGAVSWTT NDVVQNFYNPLLPSGMKFEIGQALYLGFISSSLSL IGGTLLCLSCQDEAPYRPYQAPPRATTTTANTAP AYQPPAAYKDNRAPSVTSATHSGYRLNDYV
3190		267	1037	DRMAWQGLVLAACLLMFPSTTADCLSRCSLCA VKTQDGPKPINPLICSLQCQAALLPSEEWERCQSF LSFFTPSTLGLNDKEDLGSKSVGEGPYSELAKLS GSFLKELEKSKFLPSISTKENTLSKSLEEKLRGLS DGFREGAESELMRDAQLNDGAMETGTLYLAEE DPKEQVKRYGGFLRKYPKRSSEVAGEGDGDSM GHEDLYKRYGGFLRRIRPKLKWDNQKRYGGFLR RQFKVVTRSQEDPNAYSGELFDA
3191	A	29	574	GTSAGAQTKGALCQLKVPTEKLPSPLPTMADEID FTTGDAGASSTYPMQCSALRKNGFVVLKGRPCK IVEMSTSKTGKHGHAKVHLVGIDIFTGKKYEDIC PSTHNMDVPNIKRNDYQLICIQDGYLSLLTETGE VREDLKLPEGELGKEIEGKYNAGEDVQVSVMCA MSEEYAVAIKPCK
3192	A	105	1661	KVSADGMQSCESSGDSADDPLSRGLRRRGQPRV VVIGAGLAGLAAAKALLEQGFTDVTVLEASSHIG GRVQSVKLGHATFELGATWIHGSHGNPIYHLTE ANGLLEETTDGERSVGRISLYSKNGVACYLTNH GRRIPKDVVEEFSDLYNEVYNLTQEFFRHDKPVN AESQNSVGVFTREEVRNRIRNDPDDPEATKRLKL AMIQQYLKVESCESSSHSMDEVSLSAFGEWTEIP GAHHIIPSGFMRVVELLAEGIPAHVIQLGKPVRCI HWDQASARPRGPEIEPRGEGDHNHDTGEGGQGG EEPRGGRWDEDEQWSVVVECEDCELIPADHVIV TVSLGVLKRQYTSFFRPGLPTEKVAAIHRLGIGTT DKIFLEFEEPFWGPECNSLQFVWEDEAESHTLTY PPELWYRKICGFDVLYPPERYGHVLSGWICGEEA LVMEKCDDEAVAEICTEMLRQFTGNPNIPKPRRI LRSAWGSNPYFRGSYSYTQVGSSGADVEKLAKP LPYTESSKTATK
3193	A	1	1928	QLGTRRCLRGDKVTNAMQDFLVTNLEPRFIEPQT ANLSVVFKDSNSTTPLIFVLSPGTDPAADLYKFA EEMKFSKKLSAISLGQGQGPRAEAMMRSSIERGK WVFFQNCHLAPSWMPALERLIEHINPDKVHRDF

Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, Predicted end **SEQ ID** Method Predicted nucleotide E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, beginning NO: I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, nucleotide location corresponding N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, location T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, corresponding to last amino X=Unknown, *=Stop codon, /=possible nucleotide deletion, acid residue of to first amino acid residue of peptide \=possible nucleotide insertion peptide sequence sequence RLWLTSLPSNKFPVSILQNGSKMTIEPPRGVRAN LLKSYSSLGEDFLNSCHKVMEFKSLLLSLCLFHG NALERRKFGPLGFNIPYEFTDGDLRICISQLKMFL DEYDDIPYKVLKYTAGEINYGGRVTDDWDRRCI MNILEDFYNPDVLSPEHSYSASGIYHQIPPTYDLH GYLSYIKSLPLNDMPEIFGLHDNANITFAQNETFA LLGTIIQLQPKSSSAGSQGREEIVEDVTQNILLKVP **EPINLQWVMAKYPVLYEESMNTVLVQEVIRYNR** LLQVITQTLQDLLKALKGLVVMSSQLELMAASL YNNTVPELWSAKAYPSLKPLSSWVMDLLQRLDF LQAWIQDGIPAVFWISGFFFPQAFLTGTLQNFAR KFVISIDTISFDFKVMFEAPSELTQRPQVGCYIHG LFLEGARWDPEAFQLAESQPKELYTEMAVIWLL PTPNRKAQDQDFYLCPIYKTLTRAGTLSTTGHST NYVIAVEIPTHOPORHWIKRGVALICALDY 1023 **DGWTPVHAAVDTGNVDSLKLLMYHRIPAHGNS** 3194 Â 1 FNEEESESSVFDLDGGEESPEGISKPVVPADLINH ANREGWTAAHIAASKGFKNCLEILCRHGGLEPE RRDKCNRTVHDVATDDCKHLLENLNALKIPLRIS VGEIEPSNYGSDDLECENTICALNIRKQTSWDDFS KAVSQALTNHFQAISSDGWWSLEDVTCNNTTDS NIGLSARSIRSITLGNVPWSVGQSFAQSPWDFMR KNKAEHITVLLSGPQEGCLSSVTYASMIPLQMM QNYLRLVEQYHNVIFHGPEGSLQDYIVHQLALCL KHRQMGWQDSPVEIVEELEVGCWFFPREQLLRT **CSLVA** MAASAQVSVTFEDVAVTFTQEEWGQLDAAQRT 3195 A 1 1809 LYOEVMLETCGLLMSLGCPLFKPELIYQLDHRQE LWMATKDLSQSSYPGDNTKPKTTEPTFSHLALPE EVLLQEQLTQGASKNSQLGQSKDQDGPSEMQEV HLKIGIGPORGKLLEKMSSERDGLGSDDGVCTKI TOKOVSTEGDLYECDSHGPVTDALIREEKNSYK CEECGKVFKKNALLVQHERIHTQVKPYECTECG KTFSKSTHLLQHLIIHTGEKPYKCMECGKAFNRR SHLTRHQRIHSGEKPYKCSECGKAFTHRSTFVLH HRSHTGEKPFVCKECGKAFRDRPGFIRHYIIHTGE KPYECIECIECGKAFNRRSYLTWHQQIHTGVKPF **ECNECGKAFCESADLIQHYIIHTGEKPYKCMECG** KAFNRRSHLKQHQRIHTGEKPYECSECGKAFTH CSTFVLHKRTHTGEKPYECKECGKAFSDRADLIR HFSIHTGEKPYECVECGKAFNRSSHLTRHQQIHT GEKPYECIQCGKAFCRSANLIRHSIIHTGEKPYEC SECGKAFNRGSSLTHHORIHTGRNPTIVTDVGRP **FMTAQTSVNIQELLLGKEFLNITTEENLW** 3196 1400 264 VGFWERPLRSSRWFRRSLRRWEMLARAARGTG A ALLLRGSLLASGRAPRRASSGLPRNTVVLFVPQQ **EAWVVERMGRFHRILEPGLNILIPVLDRIRYVQSL** KEIVINVPEOSAVTLDNVTLQIDGVLYLRIMDPY KASYGVEDPEYAVTQLAQTTMRSELGKLSLDKV FRERESLNASIVDAINQAADCWGIRCLRYEIKDIH **VPPRVKESMQMQVEAERRKRATVLESEGTRESA** INVAEGKKQAQILASEAEKAEQINQAAGEASAVL AKAKAKAEAIRILAAALTQHNGDAAASLTVAEQ YVSAFSKLAKDSNTILLPSNPGDVTSMVAQAMG VYGALTKAPVPGTPDSLSSGSSRDVQGTDASLDE

SEQ ID NO:	Method	Predicted beginning	Predicted end nucleotide	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine,
		nucleotide location	location corresponding	I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine,
		corresponding	to last amino	T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine,
		to first amino	acid residue of	X=Unknown, *=Stop codon, /=possible nucleotide deletion,
		acid residue of peptide	peptide sequence	=possible nucleotide insertion
		sequence		TI DRIWA (C
2107	A	66	3632	ELDRVKMS LWECAAAAAGQRDGGVTLFLKGRVLGRRCAAS
3197	A	00	3032	LFAREVCVSTSSSRPACFLHCARARGEQMHQMA
	†	-	-	SGVGSMKRSPRKMWRPGEKKEPQGVVYEDVRD
				DTEDFKEPLKVVFEGSAYGLQNFNKQKKLKTCD
				DMDTFFLHYAAAEGQIELMEKITRDSSLEVLHE
				MDDYGNTPLHCAVEKNQIESVKFLLSRGANPNL
				RNFNMMAPLHIAVQGMNNEVMKVLLEHRTIDV
		ļ		NLEGENGNTAVIIACTTNNSEALQILLNKGAKPC KSNKWGCFPIHQAAFSGSKECMEIILRFGEEHGY
				SROLHINFMNNGKATPLHLAVQNGDLEMIKMCL
				DNGAQIDPVEKGRCTAIHFAATQGATEIVKLMIS
				SYSGSVDIVNTTDGCHETMLHRASLFDHHELAD
		*		YLISVGADINKIDSEGRSPLILATASASWNIVNLL
				LSKGAQVDIKDNFGRNFLHLTVQQPYGLKNLRP
				EFMQMQQIKELVMDEDNDGCTPLHYACRQGGP GSVNNLLGFNVSIHSKSKDKKSPLHFAASYGRIN
				TCQRLLQDISDTRLLNEGDLHGMTPLHLAAKNG
				HDKVVQLLLKKGALFLSDHNGWTALHHASMGG
				YTQTMKVILDTNLKCTDRLDEDGNTALHFAARE
			·	GHAKAVALLLSHNADIVLNKQQASFLHLALHNK
				RKEVVLTIIRSKRWDECLKIFSHNSPGNKCPITEM IEYLPECMKVLLDFCMLHSTEDKSCRDYYIEYNF
				KYLQCPLEFTKKTPTQDVIYEPLTALNAMVQNN
				RIELLNHPVCKEYLLMKWLAYGFRAHMMNLGS
				YCLGLIPMTILVVNIKPGMAFNSTGIINETSDHSEI
				LDTTNSYLIKTCMILVFLSSIFGYCKEAGQIFQQK
				RNYFMDISNVLEWIIYTTGIIFVLPLFVEIPAHLQ WQCGAIAVYFYWMNFLLYLQRFENCGIFIVMLE
		l		VILKTLLRSTVVFIFLLLAFGLSFYILLNLQDPFSS
				PLLSIIQTFSMMLGDINYRESFLEPYLRNELAHPV
				LSFAQLVSFTIFVPIVLMNLLIGLAVGDIAEVQKH
				ASLKRIAMQVELHTSLEKKLPLWFLRKVDQKSTI VYPNKPRSGGMLFHIFCFLFCTGEIRQEIPNADKS
				LEMEILKQKYRLKDLTFLLEKQHELIKLIIQKMEII
				SETEDDDSHCSFQDRFKKEQMEQRNSRWNTVLR
				AVKAKTHHLEP
3198	A	51	2177	KEKSLHHVDQRPPLWHPGRPGTSQSAAMNASSE
			•	GESFAGSVQIPGGTTVLVELTPDIHICGICKQQFN NLDAFVAHKQSGCQLTGTSAAAPSTVQFVSEET
				VPATQTQTTTRTITSETQTITVSAPEFVFEHGYQT
				YLPTESNENQTATVISLPAKSRTKKPTTPPAQKRL
				NCCYPGCQFKTAYGMKDMERHLKIHTGDKPHK
			•	CEVCGKCFSRKDKLKTHMRCHTGVKPYKCKTC
				DYAAADSSSLNKHLRIHSDERPFKCQICPYASRN SSQLTVHLRSHTGDAPFQCWLCSAKFKISSDLKR
				HMRVHSGEKPFKCEFCNVRCTMKGNLKSHIRIK
[[1		HSGNNFKCPHCAFLGDSKATLRKHSRVHQSEHR
				EKCSECSYSCSSKAALRIHERIHCTVRPFKCNYCS
				FDSKQPSNLSKHMKKFHGDMVKTEALERKDTG
				RQSSRQVAKLDAKKSFHCDICDASFMREDSLRS
			·	HKRQHSEYNESKNSDVTVLQFQIDPSKQPATPLT VGHLQVPLQPSQVPQFSEGRVKIIVGHQVPQANT
				IVQAAAAAVNIVPPALVAQNPEELPGNSRLQILR
				QVSLIAPPQSSRCPSEAGAMTQPAVLLTTHEQTD
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SEQ ID NO:	Method	Predicted beginning nucleotide location corresponding to first amino acid residue of peptide sequence	Predicted end- nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion
				GATLHQTLIPTASGGPQEGSGNQTFITSSGITCTD FEGLNALIQEGTAEVTVVSDGGQNIAVATTAPPV FSSSSQQELPKQTYSIIQGAAHPALLCPADSIPD
3199	A	13	2247	QSFHSMEGDPSGLPLLARGASCYSLICPCPRPAD WSILQGTDWSILQSADWCIYNPLARHRALTGVFL QSADWCTYNPLARQKSSPSPHSTQEVQLASPLTR RPNKKDSAERNHRPAREGSVAQRQPNPAALEKA EPAARKRNEREGGGSQEPGREHSLEKGYWAPGL GPDPSMCSKQVDPSEGASSHLKHRGGSRAAHLE VRRLLRRLVGALVAEAGFCYVQVAEGQRVVGV LEVAEAAAAPVQHEPTAAVATQSRWFPRGTRPG LCSLPIAVAALLCPGSGPGAQSGLEFVERPPPSPL AVVLARWPLPPPAGRCPRDAPEARVPEKARAEG SERENNYGCGVVGGEMTTLVLDNGAYNAKIGY SHENVSVIPNCQFRSKTARLKTFTANQIDEIKDPS GLFYILPFQKGYLVNWDVQRQVWDYLFGKEMY QVDFLDTNIIITEPYFNFTSIQESMNEILFEEYQFQ AVLRVNAGALSAHRYFRDNPSELCCIIVDSGYSF THIVPYCRSKKKKEAIIRINVGGKLLTNHLKEIISY RQLHVMDETHVINQVKEDVCYVSQDFYRDMDI AKLKGEENTVMIDYVLPDFSTIKKGFCKPREEMV LSGKYKSGEQILRLANERFAVPEILFNPSDIGIQE MGIPEAIVYSIQNLPEEMQPHFFKNIVLTGGNSLF PGFRDRVYSEVRCLTPTDYDVSVVLPENPITYAW EGGKLISENDDFEDMVVTREDYEENGHSVCEEK
3200	A	3	307	FDI AVQRIRHEMNIFRLTGDLSHLAAIVILLLKIWKTR SCAGISGKSQLLFALVFTTRYLDLFTSFISLYNTS MKVWYAIHRNVFHLQCTGLWTLNLCQLCIFN
3201	A	1	469	IRHEGRGQRGKMELVQVLKRGLQQITGHGGLRG YLRVFFRTNDAKVGTLVGEDKYGNKYYEDNKQ FFGRHRWVVYTTEMNGKNTFWDVDGSMVPPE WHRWLHSMTDDPPTTKPLTARKFIWTNHKFNVT GTPEQYVPYSTTRKKIQEWIPPSTPYK
3202	A	144	840	NSSQRIMATHALEIAGLFLGGVGMVGTVAVTVM PQWRVSAFIENNIVVFENFWEGLWMNCVRQANI RMQCKIYDSLLALSPDLQAARGLMCAASVMSFL AFMMAILGMKCTRCTGDNEKVKAHILLTAGIIFII TGMVVLIPVSWVANAIIRDFYNSIVNVAQKRELG EALYLGWTTALVLIVGGALFCCVFCCNEKSSSYR YSIPSHRTTQKSYHTGKKSPSVYSRSQYV
3203	A	2	473	KYRYRRPYPVMRKICQVGPAGLAFILNISPVAHR VALCHLAGCQEQAAWYHTLQILFFLVSAYFFSCP VPEKYFPGSCDIVGHGHQIFHAFLSICTLSQLEAIL LDYQGRQEIFLQRHGPLSVHMACLSFFFLAACSA ATAALLRHKVKARLTKKDS
3204	A .	1808		PESAPLPAFISSRILPAAWRNWCSYVVTRTISCHV QNGTYLQRVLQNCPWPMSCPGSSYRTVVRPTYK VMYKIVTAREWRCCPGHSRVSCEEVAGSSASLE PMWSGSTMRRMALRPTAFSGCLNCSKVSELTER LKVLEAKMTMLTVIEQPVPPTPATPEDPAPLWGP PPAQGSPGDGGLQDQVGAWGLPGPTGPKGDAG SRGPMGMRGPPGDPLLSNTFTETNNHWPQGPTG PPGPPGPMGPPGPPGPTGVPGSPGHIGPPGPTGPK GISGHPGEKGERGLRGEPGPQGSAGQRGEPGPKG

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SEQ ID NO:	Method	Predicted beginning nucleotide location corresponding to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion
	,			DPGEKSHWGEGLHQLREALKILAERVLILETMIG LYEPELGSGAGPAGTGTPSLLRGKRGGHATNYRI VAPRSRDERG
3205	A	2810	1652	RTSTQKWQSVFNDSQEHLERFYCNPENDRMRM KYGGQEFWADLNAMNVYETTEFDQLRRLSTPPS SNVNSIYHTVWKFFCRDHFGWREYPESVIRLIEE ANSRGLKEVRFMMWNNHYILHNSFFRREIKRRP LFRSCFILLPYLQTLGGVPTQAPPPLEATSSSQIICP DGVTSANFYPETWVYMHPSQDFIQVPVSAEDKS YRIIYNLFHKTVPEFKYRILQILRVQNQFLWEKY KRKKEYMNRKMFGRDRIINERHLFHGTSQDVVD GICKHNFDPRVCGKHATMFGQGSYFAKKASYSH NFSKKSSKGVHFMFLAKVLTGRYTMGSHGMRR PPPVNPGSVTSDLYDSCVDNFFEPQIFVIFNDDQS YPYFVIQYEEVSNTVSI
3206	A		4500	CLVDSKLWKGARSVYHQLFMSSLLMDLKYKKL FAVRFAKNYERLQSDYVTDDHDREFSVADLSVQ IFTVPSLARMLITEENLMSIIIKTFMDHLRHRDAQ GRFQFERYTALQAFKFRRVQSLILDLKYVLISKPT EWSDELRQKFLEGFDAFLELLKCMQGMDPITRQ VGQHIEMEPEWEAAFTLQMKLTHVISMMQDWC ASDEKVLIEAYKKCLAVLMQCHGGYTDGEQPIT LSICGHSVETIRYCVSQEKVSIHLPVSRLLAGLHV LLSKSEVAYKFPELLPLSELSPPMLIEHPLRCLVL CAQVHAGMWRRNGFSLVNQIYYYHNVKCRRE MFDKDVVMLQTGVSMMDPNHFLMIMLSRFELY QIFSTPDYGKRFSSEITHKDVVQQNNTLIEEMLYL IIMLVGERFSPGVGQVNATDEIKREIHQLSIKPM AHSELVKSLPEDENKETGMESVIEAVAHFKKPGL TGRGMYELKPECAKEFNLYFYHFSRAEQSKAEE AQRKLKRQNREDTALPPPVLPPFCPLFASLVNILQ SDVMLCIMGTILQWAVEHNGYAWSESMLQRVL HLIGMALQEEKQHLENVTEEHVVTFTFTQKISKP GEAPKNSPSILAMLETLQNAPYLEVHKDMIRWIL KTFNAVKKMRESSPTSPVAETEGTIMEESSRDKD KAERKKAEIARLRREKIMAQMSEMQRHFIDEN KELFQQTLELDASTSAVLDHSPVASDMTLTALGP AQTQVPEQRGFVTCILCQEEQEVKVESRAMVLA AFVQRSTVLSKNRSKFIQDPEKYDPLFMHPDLSC GTHTSSCGHIMHAHCWQRYFDSVQAKEQRRQQ RLRLHTSYDVENGFFLCPLCECLSNTVIPLLLPPR NIFNNRLNFSDQPNLTQWIRTISQQIKALQFLRKE ESTPNNASTKNSENVDELQLPEGFRPDFRPKIPYS ESIKEMLTTFGTATYKVGLKVHPNEEDPRVPIMC WGSCAYTIQSIERILSDEDKPLFGPLPCRLDDCLR SLTRFAAAHWTVASVSVVQGHFCKPFASLVPND SHEELPCILDIDMFHLLVGLVLAFPALQCQDFSGI SLGTGDLHIFHLVTMAHIIQILLTSCTEENGMDQE NPPCEEESAVLALYKTLHQYTGSALKEIPSGWHL WRSVRAGIMPFLKCSALFFHYLNGVPSPPDIQVP GTSHFEHLCSYLSLPNNLICLFQENSEIMNSLIES WCRNSEVKRYLEGERDAIRYPRESNKLINLPEDY SSLINQASNFSCPKSGGDKSRAPTLCLVCGSLLCS
				QSYCCQTELEGEDVGACTAHTYSCGSGVGIFLR VRECQVLFLAGKTKGCFYSPPYLDDYGETDQGL

	,		D	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid,
SEQ ID NO:	Method	Predicted beginning nucleotide location corresponding to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion
				RRGNPLHLCKERFKKIQKLWHQHSVTEEIGHAQ EANQTLVGIDWQHL
3207	A	49	963	QLSPSQAPAGAQEVARRVTVGSASHGGRRSTMA TTVSTQRGPVYIGELPQDFLRITPTQQQRQVQLD AQAAQQLQYGGAVGTVGRLNITVVQAKLAKNY GMTRMDPYCRLRLGYAVYETPTAHNGAKNPRW NKVIHCTVPPGVDSFYLEIFDERAFSMDDRIAWT HITIPESLRQGKVEDKWYSLSGRQGDDKEGMINL VMSYALLPAAMVMPPQPVVLMPTVYQQGVGY VPITGMPAVCSPGMVPVALPPAAVNAQPRCSEE DLKAIQDMFPNMDQEVIRSVLEAQRGNKDAAIN SLLQMGEEP
3208	A	54	1196	LERTPASADMAWTKYQLFLAGLMLVTGSINTLS AKWADNFMAEGCGGSKEHSFQHPFLQAVGMFL GEFSCLAAFYLLRCRAAGQSDSSVDPQQPFNPLL FLPPALCDMTGTSLMYVALNMTSASSFQMLRGA VIIFTGLFSVAFLGRRLVLSQWLGILATIAGLVVV GLADLLSKHDSQHKLSEVITGDLLIIMAQIIVAIQ MVLEEKFVYKHNVHPLRAVGTEGLFGFVILSLLL VPMYYIPAGSFSGNPRGTLEDALDAFCQVGQQP LIAVALLGNISSIAFFNFAGISVTKELSATTRMVL DSLRTVVIWALSLALGWEAFHALQILGFLILLIGT ALYNGLHRPLLGRLSRGRPLAEESEQERLLGGTR
3209	A	104	1999	AKVVSLKEFSCFWRREKPVSSLSSLQVKAEASW DSAVHGCPQLSRGTPVDERLFLIVRVTVQLSHPA DMQLVLRKRICVNVHGRQGFAQSLLKKMSHRSS IPGCGVTFEIVSNIPEDAQGVEEREALARMAANV ENPASADSEAYIEKYLRSVLAVENLLTLDRLRQE VAVKEQLTGKGKLSRRSISSPNVNRLSGSRQDLIP SYSLGSNKGRWESQQDVSQTTVSRGIAPAPALSV SPQNNHSPDPGLSNLAASYLNPVKSFVPQMPKLL KSLFPVRDEKRGKRPSPLAHQPVPRIMVQSASPDI RVTRMEEAQPEMGPDVLVQTMGAPALKICDKP AKVPSPPPVIAVTAVTPAPEAQDGPPSPLSEASSG YFSHSVSTATLSDALGPGLDAAAPPGSMPTAPEA EPEAPISHPPPPTAVPAEEPPGPQQLVSPGRERPDL EAPAPGSPFRVRRVRASELRSFSRMLAGDPGCSP GAEGNAPAPGAGGQALASDSEEADEVPEWLREG EFVTVGAHKTGVVRYVGPADFQEGTWVGVELD LPSGKNDGSIGGKQYFRCNPGYGLLVRPSRVRR ATGPVRRSTGLRLGAPEARRSATLSGSATNLAS LTAALAKADRSHKNPENRKSWAS
3210	A	324	694	SPFWTEKRRMEKPLFPLVPLHWFGFGYTALVVS GGIVGYVKTGSVPSLAAGLLFGSLAGLGAYQLY QDPRNVWGFLAATSVTFVGVMGMRSYYYGKF MPVGLIAGASLLMAAKVGVRMLMTSD
3211	A	1078	594	VGMELPAVNLKVILLGHWLLTTWGCIVFSGSYA WANFTILALGVWAVAQRDSIDAISMFLGGLLATI FLDIVHISIFYPRVSLTDTGRFGVGMAILSLLLKPL SCCFVYHMYRERGGELLVHTGFLGSSQDRSAYQ TIDSAEAPADPFAVPEGRSQDARGY
3212	A	1	1962	FRCGLAPKGRPRRRADPVASAIMDPAEAVLQEK ALKFMMEFRSWCPGWNTMARSRLTATSTSRVQ CSMPRSLWLGCSSLADSMPSLRCLYNPGTGALT

SQ ID NO: Method Predicted beginning in incleated to be beginning in incleated to incleate the beginning in incleated to incleate the property of the pr					
AFONSSEREDCNNGEPPRKIIPEKNSLRQTYNSCA RLCLNGETVCLASTAMKTENCVAKTKLANGTSS MIVPKQRKLSASYEKEKELCVKYFEQWSESDQV EFVHEILISQMCHYQHGHINSYLKPMLQRDFITAL PARGLDHIAENILSYLDAKSLCAAELVCKEWYR VTSDGMLWKKLIERMYRTDSLWRGLAERRGWG QYLFKNKPPDGRAPPNSTYRALYPKIIQDEITES NWRCGRISLQRHICRSETSKGVYCLQYDDQKIV SGIRDNTIKIVDKNTLECKRLITGHTGSVLCLQY DERVIITGSSDSTVRVWDVNTGEMLNTLIHHCEA VLHLRRNNGMAVTCSKDRSIAVWDMASPTDITL RVILVGHRAAVNVVDFDDKYTVSASGDRTIKV WNTSTCEFVRTLNGHKRGIACLQYRDRLVVSGS SDNTIRLWDIEGGACLRVLEGHEELJVRCIRRDNIK RIVSGAYDGKIKVWDLVAALDPRAPAGTICLRT LVEHSGRVFRLOFDEFQIVSSSHDDTILLWDFLND PAAQSEPPRSPRTYTYISR 3213 A 1 1962 FRCGLAPKGRPRRRADPVASAIMDPAEAVLQEK ALKFMMEFRSWCFGWNTMARSRLTATSTSRVQ CSMPRSLWLGCSSLADSMPSLRCLYNPGTGALT AFQNSSERBCNNGPPRKIIPERNSLRQTYNSCA RLCLNQETVCLASTAMKTENCVAKTKLANGTSS MIVPKQRKLSASYBEKEKELCVKYFEQWSESDQV EFVEHLISQMCHYQHGHINSYLKPMLQRDFITAL PARGLDHIAENILSYLDAKSLCAAELVCKEWYR VTSDGMLWKKLIERMYRTDSLWRGLAERRGW QYLFKNKPPDGRAPPNSYRALYPKIQDETIES NWRCGRISLGRIHCRSETSKGVYCLQYDDGXIV SGLRDNTIKLWDLECKRLITGHTGSVLCLQY DERVITICSSDSTVRVWDWNTGEMLNTLHHICGA VLHLRFNNGMMVTCSKDRSIAVWDMASPTDITL RRVLVGHRAAVNVVDFDKYTVSASGDRTIKV WNTSTCEFVRLNGHKRGJACLQYRDRLVVSGS SDNTIRLWDIECGACLRVLEGHEELYRCIRFDNK RIVSGAYDGKIKWWDLVAALDPRAPAGTICLTL LVEHSGRVPRLQFDEFQIVSSSHDDTILIWDFLND PAQOSEPPRSPSRTYTYTISR 3214 A 1 1962 FRCGLAPKGRPRRRADPVASAIMDPAEAVLQEK ALKFMMEFRSWCPGWNTMARSRLTATSTSRVQ CSMPRSLWLGCSSLADSMPSLRCLYNPGTGALT LVEHSGRVPRLQFDEFQIVSSSHDDTILIWDFLND PAQOSEPPRSPSRTYTYTYSR SMITPKORKLISASYEKEKELCVKYPGGWSSDQV EFVEHLISQMCHYQHGHINSYLKPMLQRDFTNAL PARGLDHIAENILSYLDAKSLCAAELVCKEWYR VTSDGMLWKKLIERMYRTDSLWRGLAERRGWG QYLFKNKPPDGAPNSTYRALYPKHQDETIES NWRCGRISLQRHCRSETSKGVYCLQYDDQKIV SGLRDNTIKIWDKNTLECKRLTGHTGSVLCLQV DERVITIGSSBSTVRWDWNTGEMLNTLHHCRA VLHLRFNNGMMVTCSKDRSLAWDMASPTDITL RRVLVGHRAAVNVDFDDKYVSASGDRTIKV WNTSTCEFVRTLNGHKRGIACLQYRDRLVVSGS SDNTTIKW WDIEGGACLRVLEGHEELYRCCRFDDKK SGRDDNTIKINGHKRGIACLQYRDRLVVSGS SDNTTRLWDBECGACLRVLEGHEELYRCCRFDDKK SGRDDNTKIKWDKNATECKRLTGHTGSVLCLQU DEFVERTINGHKRGIACLQYRDRLVVSGS SDNTTRLWDBECGACLRVLEGHEELYRCCRFDDKK SGNTTRLWDBECGACLRVLEGHEELYRCCRFDDKK SONTR		Method	beginning nucleotide location corresponding to first amino acid residue of	location corresponding to last amino acid residue of peptide	I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion,
RICLNOETVCLASTAMKTENCVAKTKLANGES MIVPKORKLSASYPEKERELCVKYPFGWSESDGV EFVEHLISQMCHYQHGHINSYLKPMLQRDFITAL PARGLDHIABRILSYLDAKSLCAAELVCKEWYR VTSDGMLWKKLIERMWRITDSLWRGLAERRGG QYLFKNKPPDGNAPPNSFYRALYPKIQDIETIES NWRCGRISLGRIHCKSETSKGVYCLQYDDGKIV SGLRDNTIKIWDKNTLECKERLTGHTGSYLCLQY DERVITGSSDSTVRVWDVNTOEMLNITLHEICA VLHLRRINNGMMVTCSKDRSIAVWDMASPTDITL RRVLVGHRAAVNVVDFDDKYIVSASGDRTIKV WNTSTCEFVRTLNGHKRGIACLQYRDRLVVSGS SDNTTRLWDIECGACLRVLEGHEELVRCIRFDNK RIVSGAYDGKIKVWDLVAALDPRAPAGTLLTL LVEHSGRVFRLQFDEFQIVSSSHDDTILIWDFLND PAAQSEPPRSPSRTYTYISR 1962 FRCGLAPKGRPRRADPVASAIMDPAEAVLQEK ALKFMMEFRSWCFGWNTMARSRLTATSTSRVQ CSMPRSLWLGCSSLADSMPSLRCLYNPGTGALT AFQNSSEREDCNNGEPPRKIPEKNSLRQTYNSCA RLCLNOETVCLASTAMKTENCVAKTKLANGTSS MIVPKGRKLSASYEKEKELCVKYFEGWSESDQV EFVEHLISQMCHYQHGHINSYLKPMLQRDFITAL PARGLDHIAENILSYLDAKSLCAAELVCKEWYR VTSDGMLWKKLIERMWRTDSLWGLAERRGWG QYLFKNKPPDGNAPPNSYRALTYPKINQDETES NWRCGRHSLQRIHCRSETSKGVYCLQYDDQKIV SGLRDNTIKIWDKNTLEKRILTGHTGSVLCLQY DERVITGSSDSTVRVWDVNTGEMINITLHHICEA VLHLRRINGMWTCSKDRSIAVWDMASPTDITL RRVLVGHRAAVNVDFDDKYNSASGDRTIKV WNTSTCEFVRTLNGHKRGIACLQYRDRLVVSGS SDNTIRLWDIECGACLRVLEGHEELVRCIRFDNK RIVSGAYDGKIKVWDLVAALDPRAPAGTICLRT LVEHSGRVFRLQFDEFGQNSSHDDTILLWDFLND PAAQSEPPRSPSRTYTYISS 1962 FRCGLAPKGRPRRRADPVASAIMDPAEAVLQEK ALKFMMEFRSWCPGWNTMARSRTLATSTSTSVC CSMPRSLWLGCSSLADSMPSLRCLYNPGTGALT AFQNSSEREDCNNGEPPRKUPEKNSLRGTYNSCA RLCLNOETVCLASTAMKTENCVAKTYANGEN MIVPKGRKLSASYEKEKELCVKYFEGWSSEDQV EFVEHLISQMCHYQHGHNISYLKPMLQRSTANT PARGLEHIAERILSYLDAKSLCAAELVCKEWYR VTSDGMLWKLIERMVRTDSLWRGLAERRGWG QYLFKNKPPDGNAPPNSYRALTPATSTRYC CSMPRSLWLGCSSLADSMPSLRCLYNPGTGALT AFQNSSEREDCNNGEPPRKUPEKNSLRGTYNSCA RLCLNOETVCLASTAMKTENCVAKTKLANGTS MIVPKGRKLSASYEKEKELCVKYFEGWSSEDQV EFVEHLISQMCHYQHGHNISYLKPMLQRDFTTAL PARGLDHIAERILSYLDAKSLCAAELVCKEWYR VTSDGMLWKLIERMVRTDSLWRGLAERRGWG QYLFKNKPPDGNAPPNSYRALYPKUIQDIETIES NWRCGRISLQRHCSSETSKGVYCLQYDDQKIV SGLRDNTIKIWDKNTLECKELTGHTGGSVLCLQV DERWITGSSDSTVRVWDVNTGEMLNTLHHCEA VHLHRRPNOGMWTCSKDRSLAWDMASPTDITL RRVLVGHRAAVNVDFDDKYVSASGDRTIKV WNTSTCEFVRTLINGHKRGIACLQYRDRLVVSGS SDNTIRLWDBECGACLRVLEGHEELVRCR			sequence		
NIWRCGRHSLQRIHCRSETSKGYYCLQYDDQKIV SGLRDNTIKIWDKNTLECKRILTGHTGSVLCLQY DERVIITGSSDSTVRVWDVNTGEMLNTLIHHCEA VLHILFFNNGMMYTCSKDRSIAVWDMASFTDITL RRVLVGHRAAVNVVDFDDKYTVSASGDRTIKV WNTSTCEFVRTLNGHKRGIACLQYRDRLVVSGS SDNTIRLWDIECGACLRVLEGHEBLYRCIRFDNK RIVSGAYDGKIKVWDLVAALDPRAPAGTLCLRT LVEHSGRVFRLQFDEFQIVSSSHDDTILIWDFLND PAAQSEPPRSPSRTYTYISR 3213 A 1 1962 FRCGLAPKGRPRRADPVASAIMDPAEAVLQEK ALKFMMERRSWCPGWNTMARSRLTATSTSRVQ CSWPRSLWLGCSSLADSMPSLRCLYNPGTGALT AFQNSSEREDCNNGEPPRKIPEKNSLRQTYNSCA RLCLNQETVCLASTAMKTENCVAKTKLANGTSS MIVPKQRKLSASYBKEKELCVKYFEQWSESDQV EFVEHLISQMCHYQHGHINSYLKFMLQRDFITAL PARGLDHIAEMILSYLDAKSLCAABLVCKEWYR VTSDGMLWKKLJERMYRTDSLWRGLAERRGWG QYLFKNKPPDGNAPPNSFYRALYPKIIQDIETIES NWRCGRHSLQRIHCRSETSKGVYCLQYDDQKIV SGLRDNTIKIWDKNTLECKRILTGHTGSVLCLQY DERVITICSSDSTVRVWDVMTGEMLNTLIHECEA VLHLRFNNGMMVTCSKDRSIAVWDMASPTDITL RRVLVGHRAAVNVVDFDDKYIVSASGDRTIKV WNTSTCEFVERTLNGHKRGIACLQYRDRLVVSGS SDNTIRLWDIECGACLRVLEGHEELVRCIFFDNK RIVSGAYDGKIKVWDLVAALDPRAPAGTLCLRT LVEHSGRVFRLQFDEFQIVSSSHDDTILIWDFLND PAAQSEPPRSPSRTYTYTISR 3214 A 1 1962 FRCGLAPKGRPRRADPVASAIMDPAEAVLQEK ALKFMMERRSWCPGWNTMARSRLTATSTSRVQ CSMPRSLWLGCSSLADSMPSLRCLYNPGTGALT AFQNSSEREDCNNGEPPRKIIPEKNSLRQTYNSCA RLCLNQETVCLASTAMKTENCVAKTKLANGTSS MIVPKQRKLSASYEKERELCVKYFEGWSESDQV EFVEHLISQMCHYQHGHINSYLKPMLQRDFITAL PARGLDHABNILSYLDAKSLCAAELVCKEWYR VTSDGMLWKKLIERMVYTDSLWGLAERRGWG QYLFKNKPPDGNAPPNSFYRAL YPKIIQDIETIES NWRCGRHSLQRHCRSETSKGVYCLQYDDQKIV SGLRDNTIKIWDKNTLECKRILTGHTGSVLCLQY DERVITTGSSDSTVRVWDVNTGEMLNTLIHICEA VLHLRFNNGMMYTCSKDRSIAVWDMASFTDITL RRVLVGHRAAVNVUDFDDKYTVSASGDRTIKV WNTSTCEFVRTLNGHKRGIACLQYRDRAVLCQY GERVITTGSSDSTVRVWDVNTGBEMLNTLIHICEA VLHLRFNNGMMYTCSKDRSIAVDMASSFTDITL RRVLVGHRAAVNVUDFDDKYTVSASGDRTIKV WNTSTCEFVRTLNGHKRGIACLQYRDRAVLCQY SGLRDNTIKIWDKNTLECKRILTGHTGSVLCLQY DERVITTGSSDSTVRVWDVNTGBEMLNTLIHICEA VLHLRFNNGMMYTCSKDRSIAVWDMASSFTDITL RRVLVGHRAAVNVUDFDDKYTVSASGDRTIKV WNTSTCEFVRTLNGHKRGIACLQYRDRLVVSGS SDNTIRLWDIECGACLRVLEGHEELINGKTFDNK	e.				RLCLNQETVCLASTAMKTENCVAKTKLANGTSS MIVPKQRKLSASYEKEKELCVKYFEQWSESDQV EFVEHLISQMCHYQHGHINSYLKPMLQRDFITAL PARGLDHIAENILSYLDAKSLCAAELVCKEWYR
DERVIITGSSDSTVRVWDVNTGEMLNTLIHICEA VLHLRFNNGMMVTCSKDRSIAVWDMASPTDITIL RRVLVGHRAAVNVVDFDDKYIVSASGDRTIKV WNTSTCEFVRTLNGHKRGIACLQYRDRLVVSGS SDNTIRLWDIECGACLRVLEGHEELVRCIRFDNK RIVSGAYDGKIKVWDLVAALDPRAPAGTLCLRT LVEHSGRVFRLQFDEFQIVSSSHDDITLIWDFLND PAAQSEPPRSPSRTYTYISR 3213 A 1 1962 FRCGLAPKGRPRRADPVASAIMDPAEAVLQEK ALKFMMERRSWCPGWNTMARSRLTATSTSRVQ CSMPRSLWLGCSSLADSMPSLRCLYNPGTGALT AFQNSSEREDCNNGEPPRKIIPEKNSLRQTYNSCA RLCLNQETVCLASTAMKTENCVAKTKLANGTSS MIVPKQRKLSASYBKEKELCVKYFEQWSESDQV EFVEHLISQMCHYQHGHINSYLKPMLQRDFITAL PARGLDHIAENILSYLDAKSLCAAELVCKEWYR VTSDGMLWKKLIERMYRTDSLWRGLAERRGWG QYLFKNKPPDGNAPPNSFYRALYPKIIQDIETIES NWRCGRHSLQRHCRSETSKGVYCLQYDDQKIV SGLRDNTIKIWDKNTLECKRILTGHTGSVLCLQY DERVIITGSSDSTVRVWDVNTGEMLNTLIHHCEA VLHLRFNNGMMVTCSKDRSIAVWDMASPTDITL RRVLVGHRAAVNVVDFDDKYIVSASGDRTIKV WNTSTCEFVRTLNGHKGJACLQYRDRLVVSGG SDNTTRLWDIECGACLRVLEGHEELLVRCIRFDNK RIVSGAYDGKIKVWDLVAALDPRAPAGTLCLRT LVEHSGRVFRLQFDEFQIVSSSHDDTILIWDFLND PAAQSEPPRSPSRTYTYISR 3214 A 1 1962 FRCGLAPKGRPRRRADPVASAIMDPAEAVLQEK ALKFMMEFRSWCPGWNTMARSRLTATSTSRVQ CSMPRSLWLGCSSLADSMPSLRCLYNPGTGALT AFQNSSEREDCNNGEPPRKIIPEKNSLRQTYNSCA RLCLNQETVCLASTAMKTENCVAKTKLANGTSS MIVPKQRKLSASYEKEKELCVKYFEQWESDQV EFVEHLISQMCHYQHGHINSYLKPMLQRDFITAL PARGLDHIABRILSYLDAKSLCAAELVCKEWYA VTSDGMLWKKLIERMVRTDSLWRGLAERRGWG QYLFKNRPPDGNAPPNSFYRALYPKIIQDIETIES NWRCGRHSLGRHCRSETSRGVYCLQYDDQKIV SGLRDNTIKIWDKNTILECKRILTGHTGSVLCLQY DERVIITGSSDSTVRVWDVNTGEMLNTLIHHCES NWRCGRHSLGRHCRSETSRGVYCLQYDDQKIV SGLRDNTIKIWDKNTILECKRILTGHTGSVLCLQY DERVIITGSSDSTVRVWDVNTGEMLNTLIHHCES NWRCGRHSLGRHCRSETSRGGYYCLQYDDQKIV SGLRDNTIKIWDKNTILECKRILTGHTGSVLCLQY DERVIITGSSDSTVRVWDVNTGEMLNTLIHHCES NWRCGRHSLGRHCRSETSRGGYYCLQYDDQKIV SGLRDNTIKIWDKNTILECKRILTGHTGSVLCLQY DERVIITGSSDSTVRVWDVNTGEMLNTLIHHCES NWRCGRHSLGRINGSTSTRKGVYCLQYDDQKIV SGLRDNTIKIWDKNTILECKRILTGHTGSVLCLQY DERVIITGSSDSTVRVWDVNTGEMLNTLIHHCES SDNTIRLWDIECGACLRYLEGHEELLVRGRFDNK					NWRCGRHSLQRIHCRSETSKGVYCLQYDDQKIV
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SGLRDNTIKIWDKNTLECKRILTGHTGSVLCLQY DERVIITGSSDSTVRVWDVNTGEMLNTLIHHCEA VLHLRFNNGMMVTCSKDRSIAVWDMASPTDITL RRVLVGHRAAVNVVDFDDKYIVSASGDRTIKV WNTSTCEFVRTLNGHKRGIACLQYRDRLVVSGS SDNTIRLWDIECGACLRVLEGHEELVRCIRFDNK					
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WNTSTCEFVRTLNGHKRGIACLQYRDRLVVSGS SDNTIRLWDIECGACLRVLEGHEELVRCIRFDNK					VLHLRFNNGMMVTCSKDRSIAVWDMASPTDITL
SDNTIRLWDIECGACLRVLEGHEELVRCIRFDNK	}				

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SEQ ID NO:	Method	Predicted beginning nucleotide location corresponding to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion
				LVEHSGRVFRLQFDEFQIVSSSHDDTILIWDFLND PAAQSEPPRSPSRTYTYISR
3215	A	2	1376	EARLVGCQRGGPARPGSYSSGAETAGRAMAAN LSRNGPALQEAYVRVVTEKSPTDWALFTYEGNS NDIRVAGTGEGGLEEMVEELNSGKVMYAFCRV KDPNSGLPKFVLINWTGEGVNDVRKGACASHVS TMASFLKGAHVTINARAEEDVEPECIMEKVAKA SGANYSFHKESGRFQDVGPQAPVGSVYQKTNAV SEIKRVGKDSFWAKAEKEEENRRLEEKRRAEEA QRQLEQERRERELREAARREQRYQEQGGEASPQ RTWEQQQEVVSRNRNEQESAVHPREIFKQKERA MSTTSISSPQPGKLRSPFLQKQLTQPETHFGREPA AAISRPRADLPAEEPAPSTPPCLVQAEEEAVYEEP PEQETFYEQPPLVQQQGAGSEHIDHHIQGQGLSG QGLCARALYDYQAADDTEISFDPENLITGIEVIDE GWWRGYGPDGHFGMFPANYVELIE
3216	A	936	204	AMASTLEYSPSPLRRLVGPAAGFSRAARADLSW DPMAFFTGLWGPFTCVSRVLSHHCFSTTGSLSAI QKMTRVRVVDNSALGNSPYHRAPRCIHVYKKN GVGKVGDQILLAIKGQKKKALIVGHCMPGPRMT PRFDSNNVVLIEDNGNPVGTRIKTPIPTSLRKREG EYSKVLAIAQNFV
3217	A	,	1563	MLCALLLPSLLGATRASPTSGPQECAKGSTVW CQDLQTAARCGAVGYCQGAVWNKPTAKSLPCD VCQDIAAAAGNGLNPDATESDILALVMKTCEWL PSQESSAGCKWMVDAHSSAILSMLRGAPDSAPA QVCTALSLCEPLQRHLATLRPLSKEDTFEAVAPF MANGPLTFHPRQAPEGALCQDCVRQVSRLQEAV RSNLTLADLNIQEQCESLGPGLAVLCKNYLFQFF VPADQALRLLPPQELCRKGGFCEELGAPARLTQ VVAMDGVPSLELGLPRKQSEMQMKAGVTCEVC MNVVQKLDHWLMSNSSELMITHALERVCSVMP ASITKECIILVDTYSPSLVQLVAKITPEKVCKFIRL CGNRRRARAVHDAYAIVPSPEWDAENQGSFCNG CKRLLTVSSHNLESKSTKRDILVAFKGGCSILPLP YMIQCKHFVTQYEPVLIESLKDMMDPVAVCKKV GACHGPRTPLLGTDQCALGPSFWCRSQEAAKLC NAVQHCQKHVWKEMHLHAGEHA
3218	A	1	1563	MLCALLLPSLLGATRASPTSGPQECAKGSTVW CQDLQTAARCGAVGYCQGAVWNKPTAKSLPCD VCQDIAAAAGNGLNPDATESDILALVMKTCEWL PSQESSAGCKWMVDAHSSAILSMLRGAPDSAPA QVCTALSLCEPLQRHLATLRPLSKEDTFEAVAPF MANGPLTFHPRQAPEGALCQDCVRQVSRLQEAV RSNLTLADLNIQEQCESLGPGLAVLCKNYLFQFF VPADQALRLLPPQELCRKGGFCEELGAPARLTQ VVAMDGVPSLELGLPRKQSEMQMKAGVTCEVC MNVVQKLDHWLMSNSSELMITHALERVCSVMP ASITKECIILVDTYSPSLVQLVAKITPEKVCKFIRL CGNRRRARAVHDAYAIVPSPEWDAENQGSFCNG CKRLLTVSSHNLESKSTKRDILVAFKGGCSILPLP YMIQCKHFVTQYEPVLIESLKDMMDPVAVCKKV GACHGPRTPLLGTDQCALGPSFWCRSQEAAKLC NAVQHCQKHVWKEMHLHAGEHA
3219	A	1623	572	TSAEGWKGCTCTFKDRSKLREHLRSHTQEKVVA

SEQ ID NO:	Method	Predicted beginning nucleotide location corresponding to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion
				CPTCGGMFANNTKFLDHIRRQTSLDQQHFQCSH CSKRFATERLLRDHMRNHVNHYKCPLCDMTCPL PSSLRNHMRFRHSEDRPFKCDCCDYSCKNLIDLQ KHLDTHSEEPAYRCDFENCTFSARSLCSIKSHYR KVHEGDSEPRYKCHVCDKCFTRGNNLTVHLRK KHQFKWPSGHPRFRYKEHEDGYMRLQLVRYES VELTQQLLRQPQEGSGLGTSLNESSLQGIILETVP GEPGRKEEEEEGKGSEGTALSASQDNPSSVIHVV NQTNAQGQQEIVYYVLSEAPGEPPPVPEPPSGGI MEKLQGIAEEPEIQMV
3220	A	2760	745	SLGIPSGNTRGTGLVLDGDTSYTYHLVCMGPEAS GWGQDEPQTWPTDHRAQQGVQRQGVSYSVHA YTGQPSPRGLHSENREDEGWQVYRLGARDAHQ GRPTWALRPEDGEDKEMKTYRLDAGDADPRRL CDLERERWAVIQGQAVRKSSTVATLQGTPDHGD PRTPGPPRSTPLEENVVDREQIDFLAARQQFLSLE QANKGAPHSSPARGTPAGTTPGASQAPKAFNKP HLANGHVVPIKPQVKGVVREENKVRAVPTWAS VQVVDDPGSLASVESPGTPKETPIEREIRLAQERE ADLREQRGLRQATDHQELVEIPTRPLLTKLSLITA PRRERGRPSLYVQRDIVQETQREEDHRREGLHV GRASTPDWVSEGPQPGLRRALSSDSILSPAPDAR AADPAPEVRKVNRIPPDAYQPYLSPGTPQLEFSA FGAFGKPSSLSTAEAKAATSPKATMSPRHLSESS GKPLSTKQEASKPPRGCPQANRGVVRWEYFRLR PLRFRAPDEPQQAQVPHVWGWEVAGAPALRLQ KSQSSDLLERERESVLRREQEVAEERRNALFPEV FSPTPDENSDQNSRSSSQASGITGSYSVSESPFFSPI HLHSNVAWTVEDPVDSAPPGQRKKEQWYAGIN PSDGINSEVLEAIRVTRHKNAMAERWESRIYASE EDD
3221	A	15	478	SRVFFFFFFFPAFKMSKRGRGGSSGAKFRISLGLP VGAVINCADNTGAKNLYIISVKGIKGRLNRLPAA GVGDMVMATVKKGKPELRKKVHPAVVIRQRKS YRRKDGVFLYFEDNAGVIVNNKGEMKGSAITGP VAKECADLWPRIASNAGSIA
3222	A	207	1321	PLIPLHPANRSPATMAELQEVQITEEKPLLPGQTP EAAKTHSVETPYGSVTFTVYGTPKPKRPAILTYH DVGLNYKSCFQPLFQFEDMQEIIQNFVRVHVDAP GMEEGAPVFPLGYQYPSLDQLADMIPCVLQYLN FSTIIGVGVGAGAYILARYALNHPDTVEGLVLINI DPNAKGWMDWAAHKLTGLTSSIPEMILGHLFSQ EELSGNSELIQKYRNIITHAPNLDNIELYWNSYNN RRDLNFERGGDITLRCPVMLVVGDQAPHEDAVV ECNSKLDPTQTSFLKMADSGGQPQLTQPGKLTE AFKYFLQGMGYMASSCMTRLSRSRTASLTSAAS VDGNRSRSRTLSQSSESGTLSSGPPGHTMEVSC
3223	A	132	1664	SARRWGAAGAGPHGLHLRAHGPRPSVRTGLPSV GRQAAGAAMGRGWGFLFGLLGAVWLLSSGHGE EQPPETAAQRCFCQVSGYLDDCTCDVETIDRFNN YRLFPRLQKLLESDYFRYYKVNLKRPCPFWNDIS QCGRRDCAVKPCQSDEVPDGIKSASYKYSEEAN NLIEECEQAERLGAVDESLSEETQKAVLQWTKH DDSSDNFCEADDIQSPEAEYVDLLLNPERYTGYK GPDAWKIWNVIYEENCFKPQTIKRPLNPLASGQG

SEQ ID NO:	Method	Predicted beginning nucleotide location corresponding to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \(\rightarrow \text{possible nucleotide insertion} \)
				TSEENTFYSWLEGLCVEKRAFYRLISGLHASINV HLSARYLLQETWLEKKWGHNITEFQQRFDGILTE GEGPRRLKNLYFLYLIELRALSKVLPFFERPDFQL FTGNKIQDEENKMLLLEILHEIKSFPLHFDENSFF AGDKKEAHKLKEDFRLHFRNISRIMDCVGCFKC RLWGKLQTQGLGTALKILFSEKLIANMPESGPSY EFHLTRQEIVSLFNAFGRISYKCERIRKTSRNLLQ NIH
3224	A	2	803	PGSTISWDRDAAGESGTRAASPSPSGSRTAGRLP SPSYSPLPAPSLFPPPPLPAPAASTMSAGGDFGNP LRKFKLVFLGEQSVGKTSLITRFMYDSFDNTYQA TIGIDFLSKTMYLEDRTVRLQLWDTAGQERFRSL IPSYIRDSTVAVVVYDITNLNSFQQTSKWIDDVRT ERGSDVIIMLVGNKTDLADKRQITIEEGEQRAKE LSVMFIETSAKTGYNVKQLFRRVASALPGMENV QEKSKEGMIDIKLDKPQEPPASEGGCSC
3225	A	3	5054	PEVTKPSLSQPTAASPIGSSPSPPVNGGNNAKRVA VPNGQPPSAARYMPREVPPRFRCQQDHKVLLKR GQPPPPSCMLLGGGAGPPPCTAPGANPNNAQVT GALLQSESGTAPDSTLGGAAASNYANSTWGSGA SSNNGTSPNPIHIWDKVIVDGSDMEEWPCIASKD TESSSENTTDNNSASNPGSEKSTLPGSTTSNKGK GSQCQSASSGNECNLGVWKSDPKAKSVQSSNST TENNNGLGNWRNVSGQDRIGPGSGFSNFNPNSN PSAWPALVQEGTSRKGALETDNSNSSAQVSTVG QTSREQQSKMENAGVNFVSGREQAQIHNTDGP KNGNTNSLNLSSPNPMENKGMPFGMGLGNTSRS TDAPSQSTGDRKTGSVGSWGAARGPSGTDTVSG QSNSGNNGNNGKEREDSWKGASVQKSTGSKND SWDNNNRSTGGSWNFGPQDSNDNKWGEGNKM TSGVSQGEWKQPTGSDELKIGEWSGPNQPNSST GAWDNQKGHPLLENQGNAQAPCWGRSSSSTGS EVEGQSTGSNHKAGSSDSHNSGRRSYRPTHPDC QAVLQTLLSRTDLDPRVLSNTGWGQTQIKQDTV WDIEEVPRPEGKSDKGTEGWESAATQTKNSGG WGDAPSQSNQMKSGWGELSASTEWKDPKNTGG WNDYKNNNSSNWGGGRPDEKTPSSWNENPSKD QGWGGGRQPNQGWSSGKNGWGEEVDQTKNSN WESSASKPVSGWGEGGQNEIGTWGNGGNASLA SKGGWEDCKRSPAWNETGRQPNSWNKQHQQQ QPPQQPPPPPPEASAGSWGGPPPPPPGNVRPSNSS WSSGPQPATPKDEEPSGWEEPSPQSISRKMDIDD GTSAWGDPNSYNYKNVNLWDKNSQGGPAPREP NLPTPMTSKSASDSKSMQDGWGESDGPVTGARH PSWEEEEDGGVWNTTGSQGSASSHNSASWGQG GKKQMKCSLKGGNNDSWMNPLAKQFSNMGLL SQTEDNPSSKMDLSVGSLSDKKFDVDKRAMNLG DFNDIMRKDRSGFRPPNSKDMGTTDSGPYFEKG GSHGLFGNSTAQSRGLHTPVQPLNSSPSLRAQVP PQFISPQVSASMLKQFPNSGLSPGLFNVGPQLSPQ QIAMLSQLPQIPQFQLACQLLLQQQQQQQLLQN QRKISQAVRQQQEQQLARMVSALQQQQQQQQR QPGMKHSPSHPVGPKPHLDNMVPNALNVGLPDL QTKGPIPGYGSGFSSGGMDYGMVGGKEAGTESR FKQWTSMMEGLPSVATQEANMHKNGAIVAPGK

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SEO ID	Method	Predicted	Predicted end	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid,
NO:	Method	beginning nucleotide location	nucleotide location corresponding	E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Prolline, Q=Glutamine, R=Arginine, S=Serine,
		corresponding	to last amino	T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine,
	Ì	to first amino acid residue of	acid residue of peptide	X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion
		peptide sequence	sequence	
				TRGGSPYNQFDIIPGDTLGGHTGPAGDSWLPAKS PPTNKIGSKSSNASWPPEFQPGVPWKGIQNIDPES
				DPYVTPGSVLGGTATSPIVDTDHQLLRDNTTGSN
				SSLNTSLPSPGAWPYSASDNSFTNVHSTSAKFPD
		ļ		YKSTWSPDPIGHNPTHLSNKMWKNHISSRNTTPL
				PRPPPGLTNPKPSSPWSSTAPRSVRGWGTQDSRL
	}	i		ASASTWSDGGSVRPSYWLVLHNLTPQIDGSTLRT
				ICMQHGPLLTFHLNLTQGTALIRYSTKQEAAKAQ
				TALHMCVLGNTTILAEFATDDEVSRFLAQAQPPT PAATPSAPAAGWQSLETGQNQSDPVGPALNLFG
				GSTGLGQWSSSAGGSSGADLAGASLWGPPNYSS
				SLWGVPTVEDPHRMGSPAPLLPGDLLGGGSDSI
3226	Α	200	1387	VPWKRQDEQLSLQVETLYLDSPAVIHLLSPTFLP
				PSSLPPFLQIVDSSSSACTLDSFFPFLAPWDSPQDC
				GFKDHQPLTLQALTVELARWTLMLLLSTAMYG AHAPLLALCHVDGRVPFRPSSAVLLTELTKLLLC
				AFSLLVGWQAWPQGPPPWRQAAPFALSALLYG
	İ			ANNNLVIYLQRYMDPSTYQVLSNLKIGSTAVLY
		1		CLCLRHRLSVRQGLALLLLMAAGACYAAGGLQ
			ļ	VPGNTLPSPPPAAAASPMPLHITPLGLLLLILYCLI
				SGLSSVYTELLMKRQRLPLALQNLFLYTFGVLLN
				LGLHAGGGSGPGLLEGFSGWAALVVLSQALNGL LMSAVMKHGSSITRLFVVSCSLVVNAVLSAVLL
				RLQLTAAFFLATLLIGLAMRLYYGSR
3227	A	1	679	RSTRARTRRPGLRAVPLPVGGFLGKMKWVWAL
		1		LLLAALGSGRAERDCRVSSFRVKENFDKARFSGT
				WYAMAKKDPEGLFLQDNIVAEFSVDETGQMSA
			-	TAKGRVRLLNNWDVCADMVGTFTDTEDPAKFK MKYWGVASFLQKGNDDHWIVDTDYDTYAVQY
	}		}	SCRLLNLDGTCADSYSFVFSRDPNGLPPEAQKIV
				RQRQEELCLARQYRLIVHNGYCDGRSERNLL
3228	Α	430	1104	QQESPAAGAARMNCKEGTDSSCGCRGNDEKKM
				LKCVVVGDGAVGKTCLLMSYANDAFPEEYVPT
		İ		VFDHYAVTVTVGGKQHLLGLYDTAGQEDYNQL RPLSYPNTDVFLICFSVVNPASYHNVQEEWVPEL
		1		KDCMPHVPYVLIGTQIDLRDDPKTLARLLYMKE
				KPLTYEHGVKLAKAIGAQCYLECSALTQKGLKA
				VFDEAILTIFHPKKKKKRCSEGHSCCSII
3229	A	25	722	AISAGRSAKMQLKPMEINPEMLNKVLSRLGVAG
			1	QWRFVDVLGLEEESLGSVPAPACALLLLFPLTAQ
				HENFRKKQIEELKGQEVSPKVYFMKQTIGNSCGT IGLIHAVANNQDKLGFEDGSVLKQFLSETEKMSP
		ł		EDRAKCFEKNEAIQAAHDAVAQEGQCRVDDKV
		·		NFHFILFNNVDGHLYELDGRMPFPVNHGASSEDT
				LLKDAAKVCREFTEREQGEVRFSAVALCKAA
3230	A	282	1479	GDAATTACAPPDWFLGPRKLAAGPAGGGMLPR
			,	RLLAAWLAGTRGGGLLALLANQCRFVTGLRVR RAQQIAQLYGRLYSESSRRVLLGRLWRRLHGRP
				GHASALMAALAGVFVWDEERIQEEELQRSINEM
1		1		KRLEEMSNMFQSSGVQHHPPEPKAQTEGNEDSE
				GKEQRWEMVMDKKHFKLWRRPITGTHLYQYRV
				FGTYTDVTPRQFFNVQLDTEYRKKWDALVIKLE
				VIERDVVSGSEVLHWVTHFPYPMYSRDYVYVRR
1		1		YSVDQENNMMVLVSRAVEHPSVPESPEFVRVRS YESQMVIRPHKSFDENGFDYLLTYSDNPQTVFPR
	1	1	L	TENGRIAM THEOLDEHOLD LPPT LODINLALLY

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SEQ ID NO:	Method	Predicted beginning nucleotide location corresponding to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion
				YCVSWMVSSGMPDFLEKLHMATLKAKNMEIKV KDYISAKPLEMSSEAKATSQSSERKNEGSCGPAR IEYA
3231	A	2117	590	FVPEPPEAGASSPCAPGDPDMSFRKVVRQSKFRH VFGQPVKNDQCYEDIRVSRVTWDSTFCAVNPKF LAVIVEASGGGAFLVLPLSKTGRIDKAYPTVCGH TGPVLDIDWCPHNDEVIASGSEDCTVMVWQIPE NGLTSPLTEPVVVLEGHTKRVGIIAWHPTARNVL LSAGCDNVVLIWNVGTAEELYRLDSLHPDLIYN VSWNHNGSLFCSACKDKSVRIIDPRRGTLVAERE KAHEGARPMRAIFLADGKVFTTGFSRMSERQLA LWDPENLEEPMALQELDSSNGALLPFYDPDTSV VYVCGKGDSSIRYFEITEEPPYIHFLNTFTSKEPQR GMGSMPKRGLEVSKCEIARFYKLHERKCEPIVM TVPRKSDLFQDDLYPDTAGPEAALEAEEWVSGR DADPILISLREAYVPSKQRDLKISRRNVLSDSRPA MAPGSSHLGAPASTTTAADATPSGSLARAGEAG KLEEVMQELRALRALVKEQGDRICRLEEQLGRM ENGDA
3232	A	3	718	RLREDDRRGLPLSSPLWTEPPLSCCLPATYPADM GTAGAMQLCWVILGFLLFRGHNSQPTMTQTSSS QGGLGGLSLTTEPVSSNPGYIPSSEANRPSHLSST GTPGAGVPSSGRDGGTSRDTFQTVPPNSTTMSLS MREDATILPSPTSETVLTVAAFGVISFIVILVVVVI ILVGVVSLRFKCRKSKESEDPQKPGSSGLSESCST ANGEKDSITLISMKNINMNNGKQSLSAEKVL
3233	A	3	718	RLREDDRRGLPLSSPLWTEPPLSCCLPATYPADM GTAGAMQLCWVILGFLLFRGHNSQPTMTQTSSS QGGLGGLSLTTEPVSSNPGYIPSSEANRPSHLSST GTPGAGVPSSGRDGGTSRDTFQTVPPNSTTMSLS MREDATILPSPTSETVLTVAAFGVISFIVILVVVVI ILVGVVSLRFKCRKSKESEDPQKPGSSGLSESCST ANGEKDSITLISMKNINMNNGKQSLSAEKVL
3234	A	1169	4292	AGDCGRLGVGGSEFPWEGSALGASPLPPICLQSR TWLLRAPAPAELGELEEVAAGRGDVWEPFLDSP GREESLQEASPRLADHGSSSGGGWEVKRSQRLR RGPSSPRRPYQDMEYERRGGRGDRTGRYGATDR SQDDGGENRSRDHDYRDMDYRSYPREYGSQEG KHDYDDSSEEQSAEDSYEASPGSETQRRRRRRH RHSPTGPPGFPRDGDYRDQDYRTEQGEEEEEED EEEEEKASNIVMLRMLPQAATEDDIRGQLQSHG VQAREVRLMRNKSSGQSRGFAFVEFSHLQDATR WMEANQHSLNILGQKVSMHYSDPKPKINEDWL CNKCGVQNFKRREKCFKCGVPKSEAEQKLPLGT RLDQQTLPLGGRELSQGLLPLPQPYQAQGVLAS QALSQGSEPSSENANDTIILRNLNPHSTMDSILGA LAPYAVLSSSNVRVIKDKQTQLNRGFAFIQLSTIE AAQLLQILQALHPPLTIDGKTINVEFAKGSKRDM ASNEGSRISAASVASTAIAAAQWAISQASQGGEG TWATSEEPPVDYSYYQQDEGYGNSQGTESSLYA HGYLKGTKGPGITGTKGDPTGAGPEASLEPGADS VSMQAFSRPQPGAAPGIYQQSAEASSSQGTAANS QSYTIMSPAVLKSELQSPTHPSSALPPATSPTAQE SYSQYPVPDVSTYQYDETSGYYYDPQTGLYYDP NSQYYYNAQSQQYLYWDGERRTYVPALEQSAD

Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, Predicted end Predicted Method SEQ ID E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, NO: beginning nucleotide I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, location nucleotide N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, corresponding location T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, to last amino corresponding acid residue of X=Unknown, *=Stop codon, /=possible nucleotide deletion, to first amino =possible nucleotide insertion peptide acid residue of sequence peptide sequence **GHKETGAPSKEGKEKKEKHKTKTAQQIAKDME** RWARSLNKQKENFKNSFQPISSLRDDERRESATA DAGYAILEKKGALAERQHTSMDLPKLASDDRPS **PPRGLVAAYSGESDSEEEQERGGPEREEKLTDW QKLACLLCRRQFPSKEALIRHQQLSGLHKQNLEI** HRRAHLSENELEALEKNDMEQMKYRDRAAERR **EKYGIPEPPEPKRRKYGGISTASVDFEQPTRDGLG** SDNIGSRMLQAMGWKEGSGLGRKKQGIVTPIEA **OTRVRGSGLGARGSSYGVTSTESYKETLHKTMV** TRFNEAO **PSFLNTGLGPTALGVLGGAGAGLMSNPSPQVPEE** 3235 Ā 3 1217 EASTSVCRPKSSMASTSRRQRRERRFRRYLSAGR LVRAQALLQRHPGLDVDAGQPPPLHRACARHD **APALCLLIRLGADPAHQDRHGDTALHAAARQG** PDAYTDFFLPLLSRCPSAMGIKNKDGETPGOILG WGPPWDSAEEEEEDDASKEREWRQKLQGELED **EWQEVMGRFEGDASHETQEPESFSAWSDRLARE** HAQKCQQQREAEGSCRPPRAEGSSQSWRQQEE **EQRLFRERARAKEEELRESRARRAQEALGDREP** KPTRAGPREEHPRGAGRGSLWRFGDVPWPCPGG GDPEAMAAALVARGPPLEEQGALRRYLRVQQV RWHPDRFLQRFRSQIETWELGRVMGAVTALSQA LNRHAEALK **GPASGMAEPTSDFETPIGWHASPELTPTLGPLSDT** 1416 3236 A APPRDRWMFWAMLPPPPPPLTSSLPAAGSKPSSE SOPPMEAOSLPGAPPPFDAQILPGAQPPFDAQSPL DSQPQPSGQPWNFHASTSWYWRQSSDRFPRHQK SLNPAVKNSYYPRKYDAKFTDFSLPPSRKQKKK KRKEPVFHFFCDTCDRGFKNQEKYDKHMSEHTK **CPELDCSFTAHEKIVQFHWRNMHAPGMKKIKLD TPEEIARWREERRKNYPTLANIERKKKLKLEKEK** RGAVLTTTQYGKMKGMSRHSQMAKIRSPGKNH KWKNDNSRQRAVTGSGSHLCDLKLEGPPEANA DPLGVLINSDSESDKEEKPQHSVIPKEVTPALCSL **MSSYGSLSGSESEPEETPIKTEADVLAENQVLDSS** APKSPSQDVKATVRNFSEAKSENRKKSFEKTNPK REKRLSQLSNVIRTKNTPSISLGNASSSGHST 2204 **FVGEQEGGCEAGAGRGAQTYPGEAGERWFGRR** 3237 A 3806 RRRGRVVSRKKMSLKSERRGIHVDQSDLLCKKG CGYYGNPAWQGFCSKCWREEYHKARQKQIQED WELAERLOREEEEAFASSOSSOGAOSLTFSKFEE KKTNEKTRKVTTVKKFFSASSRVGSKKEIQEAKA **PSPSINROTSIETDRVSKEFIEFLKTFHKTGQEIYK OTKLFLEGMHYKRDLSIEEQSECAQDFYHNVAE** RMQTRGKVPPERVEKIMDQIEKYIMTRLYKYVF CPETTDDEKKDLAIQKRIRALRWVTPQMLCVPV NEDIPEVSDMVVKAITDIIEMDSKRVPRDKLACIT KCSKHIFNAIKITKNEPASADDFLPTLIYIVLKGNP PRLQSNIQYITRFCNPSRLMTGEDGYYFTNLCCA VAFIEKLDAQSLNLSQEDFDRYMSGQTSPRKQEA ESWSPDACLGVKQMYKNLDLLSQLNERQERIMN **EAKKLEKDLIDWTDGIAREVQDIVEKYPLEIKPP** NQPLAAIDSENVENDKLPPPLQPQVYAG VLSVCPTGVFRPAPCRMAFMKKYLLPILGLFMA 449 3238 1373 Α YYYYSANEEFRPEMLQGKKVIVTGASKGIGREM

SEQ ID	Method	Predicted	Predicted end	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid,
NO:		beginning nucleotide	nucleotide location	E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine,
		location corresponding	corresponding t last amino	N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine,
		to first amino	acid residue of	X=Unknown, *=Stop codon, /=possible nucleotide deletion,
		acid residue of peptide	peptide sequence	\=possible nucleotide insertion
ł		sequence	sequence	
				AYHLAKMGAHVVVTARSKETLQKVVSHCLELG
İ				AASAHYIAGTMEDMTFAEQFVAQAGKLMGGLD MLILNHITNTSLNLFHDDIHHVRKSMEVNFLSYV
				VLTVAALPMLKQSNGSIVVVSSLAGKVAYPMVA
		İ		AYSASKFALDGFFSSIRKEYSVSRVNVSITLCVLG
				LIDTETAMKAVSGIVHMQAAPKEECALEIIKGGA
}				LRQEEVYYDSSLWTTLLIRNPCRKILEFLYSTSYN MDRFINK
3239	A	213	422	ERTMQLEIKVALNFIIFYLYNKLLW/QPLKKK*EA
				HWYPDKPLKGSGFHT/GEMVDPVGELAAKRSGL
				TVED
3240	A	1255	1425	HESYHVNPNLCNPVAPTSGAHSIG*KWPSWLGA VAHSCNPSTLVGRGGRITRGQELR
3241	Α	161	547	PAGIGRSTAKTPGTPGSLEMENLKSGVYPLKEAS GCPGADRNLLVYSFYEKGPLTFRDVAIEFSLEEW
				GCPGADRNLLVYSFYEKGPLIFRDVAIEFSLEEW QCLDTAQQDLYRKVMLENYRNLVFLAGIAVSKP
				DLITCLEQGKEPWNMKRHAMVDQPPGR
3242	A	50	241	PLPARGKSTLPATFCSPSAPELASMSVVPPNRSQT
			,	GWPRGVTQFGNKYIQQTKPLTLERTINL
3243	Α	380	702	FVAYLKLPFFSQVCLFASSEMFFTISRKNMSQKLS LLLLVFGLIWGLMLLHYTFQQPRHQSSVKLREQI
				LDLSKRYVKALAEENKNTVDVENGASMAGYGK
				ITVEYF
3244	Α	37	1391	VLMDGRMMRSMRLREEESPGPSHTASCLCGSAP
ĺ				CILCSCCPASRNSTVSRLIFTFFLFLGVLVSIIMLSP GVESQLYKLPWVCEEGAGIPTVLQGHIDCGSLLG
				YRAVYRMCFATAAFFFFTLLMLCVSSSRDPRA
		i		AIQNGFWFFKFLILVGLTVGAFYIPDGSFTNIWFY
				FGVVGSFLFILIQLVLLIDFAHSWNQRWLGKAEE
			J	CDSRAWYAGLFFFTLLFYLLSIAAVALMFMYYT EPSGCHEGKVFISLNLTFCVCVSIAAVLPKVQDA
				QPNSGLLQASVITLYTMFVTWSALSSIPEQKCNP
				HLPTQLGNETVVAGPEGYETQWWDAPSIVGLIIF
				LLCTLFISLRSSDHRQVNSLMQTEECPPMLDATQ
				QQQQVAACEGRAFDNEQDGVTYSYSFFHFCLVL ASLHVMMTLTNWYKPGETRKMISTWTAVWVKI
				CASWAGLLLYL
3245	A	52	426	SSLGNEDDEILSLAKDITGMFVASHRKMRAHQV
		1		LTFLLLFVITSVASENASTSRGCGLDLLPQYVSLC
			1	DLDAIWGIVVEAAAGAGALITLLLMLILLVRLPF FKEKEKKSPVGLHFLFLLGTLGP
3246	A	3	515	HEVCGSGCCCHCCAGGPVARQKALPRLRGVMS
•				RFLNVLRSWLVMVSIIAMGNTLQSFRDHTFLYEK
		1		LYTGKPNLVNGLQARTFGIWTLLSSVIRCLCAIDI
	1			HNKTLYHITLWTFLLALGHFLSELFVYGTAAPTI
				GVLAPLMVASFSILGMLVGLRYLEVEPVSRQKK RN
3247	A	1	932	ERLCFPCMQSKIYSYMSPNKCSGMRFPLQEENSV
		1		THHEVKCQGKPLAGIYRKREEKRNAGNAVRSA
				MKSEEQKIKDARKGPLVPFPNQKSEAAEPPKTPP
		1		SSCDSTNAAIAKQALKKPIKGKQAPRKKAQGKT QQNRKLTDFYPVRRSSRKSKAELQSEERKRIDELI
	1			ESGKEEGMKIDLIDGKGRGVIATKQFSRGDFVVE
		1		YHGDLIEITDAKKREALYAQDPSTGCYMYYFQY
				LSKTYCVDATRETNRLGRLINHSKCGNCQTKLH

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SEQ ID NO:	Method	Predicted beginning nucleotide location corresponding to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion
				DIDGVPHLILIASRDIAAGEELLYDYGDRSKASIE AHPWLKH
3248	A	3	870	PGSTISCSELKGTQCRATAGSRGRRPPMTCWLRG VTATFGRPAEWPGYLSHLCGRSAAMDLGPMRK SYRGDREAFEETHLTSLDPVKQFAAWFEEAVQC PDIGEANAMCLATCTRDGKPSARMLLKGFGKD GFRFFTNFESRKGKELDSNPFASLVFYWEPLNRQ VRVEGPVKKLPEEEAECYFHSRPKSSQIGAVVSH QSSVIPDREYLRKKNEELEQLYQDQEVPKPKSW GGYVLYPQVMEFWQGQTNRLHDRIVFRRGLPTG DSPLGPMTHRGEEDWLYERLAP
3249	A	43	1210	TRVGRGESGLKMEVKPPPGRPQPDSGRRRRRRG EEGHDPKEPEQLRKLFIGGLSFETTDDSLREHFEK WGTLTDCVVMRDPQTKRSRGFGFVTYSCVEEV DAAMCARPHKVDGRVVEPKRAVSREDSVKPGA HLTVKKIFVGGIKEDTEEYNLRDYFEKYGKIETIE VMEDRQSGKKRGFAFVTFDDHDTVDKIVVQKY HTINGHNCEVKKALSKQEMQSAGSQRGRGGGS GNFMGRGGNFGGGGGYNGFGGRGGYGG GGGSRGSYGGGDGGYNGFGGDGGNYGGGPG YSSRGGYGGGPGYGNQGGGYGGGGYDGYN EGGNFGGGNYGGGGNYNDFGNYSGQQQSNYGP MKGGSFGGRSSGSPYGGGYGSGGGSGGYGSRRF
3250	A	32	1175	VAGRGDMAALRDAEIQKDVQTYYGQVLKRSAD LQTNGCVTTARPVPKHIREALQNVHEEVALRYY GCGLVIPEHLENCWILDLGSGSGRDCYVLSQLVG EKGHVTGIDMTKGQVEVAEKYLDYHMEKYGFQ ASNVTFIHGYIEKLGEAGIKNESHDIVVSNCVINL VPDKQQVLQEAYRVLKHGGELYFSDVYTSLELP EEIRTHKVLWGECLGGALYWKELAVLAQKIGFC PPRLVTANLITIQNKELERVIGDCRFVSATFRLFK HSKTGPTKRCQVIYNGGITGHEKELMFDANFTFK EGEIVEVDEETAAILKNSRFAQDFLIRPIGEKLPTS GGCSALELKDIITDPFKLAEESDSMKSRCVPDAA GGCCGTKKSC
3251	A	32	1175	VAGRGDMAALRDAEIQKDVQTYYGQVLKRSAD LQTNGCVTTARPVPKHIREALQNVHEEVALRYY GCGLVIPEHLENCWILDLGSGSGRDCYVLSQLVG EKGHVTGIDMTKGQVEVAEKYLDYHMEKYGFQ ASNVTFIHGYIEKLGEAGIKNESHDIVVSNCVINL VPDKQQVLQEAYRVLKHGGELYFSDVYTSLELP EEIRTHKVLWGECLGGALYWKELAVLAQKIGFC PPRLVTANLITIQNKELERVIGDCRFVSATFRLFK HSKTGPTKRCQVIYNGGITGHEKELMFDANFTFK EGEIVEVDEETAAILKNSRFAQDFLIRPIGEKLPTS GGCSALELKDIITDPFKLAEESDSMKSRCVPDAA GGCCGTKKSC
3252	A	1	574	PLGSNTAPALRVMVQAWYMDDAPGDPRQPHRP DPGRPVGLEQLRRLGVLYWKLDADKYENDPELE KIRRERNYSWMDIITICKDKLPNYEEKIKMFYEE HLHLDDEIRYILDGSGYFDVRDKEDQWIRIFMEK GDMVTLPAGIYHRFTVDEKNYTKAMRLFVGEPV WTAYNRPADHFEARGQYVKFLAQTA
3253	Α	2	984	ARAAAHCGICRLVRWWRKRRSVMGIQTSPVLLA SLGVGLVTLLGLAVGSYLVRRSRRPQVTLLDPNE

Predicted Predicted end Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, SEO ID Method E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, nucleotide NO: beginning location I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, nucleotide N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, corresponding location corresponding T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, to last amino X=Unknown, *=Stop codon, /=possible nucleotide deletion, to first amino acid residue of \=possible nucleotide insertion acid residue of peptide peptide sequence sequence KYLLRLLDKTTVSHNTKRFRFALPTAHHTLGLPV GKHIYLSTRIDGSLVIRPYTPVTSDEDQGYVDLVI KVYLKGVHPKFPEGGKMSQYLDSLKVGDVVEF RGPSGLLTYTGKGHFNIQPNKKSPPEPRVAKKLG MIAGGTGITPMLQLIRAILKVPEDPTQCFLLFANQ TEKDIILREDLEELQARYPNRFKLWFTLDHPPKD WAYSKGFVTADMIREHLPAPGDDVLVLLCGPPP MVQLACHPNLDKLGYSQKMRFTY LQSAGEGVTHVLILLESPARPVAAVTQVQRRRY 968 3254 HRLSDMSMLAERRRKQKWAVDPQNTAWSNDD SKFGQRMLEKMGWSKGKGLGAQEQGATDHIKV QVKNNHLGLGATINNEDNWIAHQDDFNQLLAEL NTCHGQETTDSSDKKEKKSFSLEEKSKISKNRVH YMKFTKGKDLSSRSKTDLDCIFGKRQSKKTPEG DASPSTPEENETTTTSAFTIQEYFAKRMAALKNK PQVPVPGSDISETQVERKRGKKRNKEATGKDVE SYLOPKAKRHTEGKPERAEAQERVAKKKSAPAE **EQLRGPCWDQSSKASAQDAGDHVQPA** 3255 173 439 **GSAAMKVKIKCWNGVATWLWVANDENCGICR** A MAFNGCCPDCKVPGDDCPLVWGQCSHCFHMHC ILKWLHAQQVQQHCPMCRQEWKFKE TAARRRQKGTAARRRQKGTLEEVVLPPRSCRVF 3256 A 2 377 WIHSGTTMSKVSFKITLTSDPRLPYKVLSVPESTP FTAVLKFAAEEFKVPAATSAIITNDGIGINPAQTA **GNVFLKHGSELRIIPRDRVGSC** GCSAAAAGAGSGPWAAQEKQFPPALLSFFIYNPR 3257 1454 A 3 **FGPREGOEENKILFYHPNEVEKNEKIRNVGLCEAI** VOFTRTFSPSKPAKSLHTQKNRQFFNEPEENFWM **VMVVRNPIIEKQSKDGKPVIEYQEEELLDKVYSS** VLRQCYSMYKLFNGTFLKAMEDGGVKLLKERL **EKFFHRYLQTLHLQSCDLLDIFGGISFFPLDKMTY** LKIQSFINRMEESLNIVKYTAFLYNDQLIWSGLEQ DDMRILYKYLTTSLFPRHIEPELAGRDSPIRAEMP GNLQHYGRFLTGPLNLNDPDAKCRFPKIFVNTD DTYEELHLIVYKAMSAAVCFMIDASVHPTLDFC RRLDSIVGPQLTVLASDICEQFNINKRMSGSEKEP OFKFIYFNHMNLAEKSTVHMRKTPSVSLTSVHPD **LMKILGDINSDFTRVDEDEEIIVKAMSDYWVVG** KKSDRRELYVILNQKNANLIEVNEEVKKLCATQF NNIFFLD **APRGCSMPHRKKKPFIEKKKAVSFHLVHRSQRD** 3258 113 1558 A PLAADESAPQRVLLPTQKIDNEERRAEQRKYGVF **FDDDYDYLOHLKEPSGPSELIPSSTFSAHNRREEK EETLVIPSTGIKLPSSVFASEFEEDVGLLNKAAPV** SGPRLDFDPDIVAALDDDFDFDDPDNLLEDDFIL OANKATGEEEGMDIQKSENEDDSEWEDVDDEK **GDSNDDYDSAGLLSDEDCMSVPGKTHRAIADHL FWSEETKSRFTEYSMTSSVMRRNEQLTLHDERFE** KFYEQYDDDEIGALDNAELEGSIQVDSNRLQEVL NDYYKEKAENCVKLNTLEPLEDQDLPMNELDES **EEEEMITVVLEEAKEKWDCESICSTYSNLYNHPQ** LIKYQPKPKQIRISSKTGIPLNVLPKKGLTAKQTE RIOMINGSDLPKVSTQPRSKNESKEDKRARKQAI KEERKERRVEKKANKLAFKLEKRROEKELLNLK KNVEGLKL

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SEQ ID NO:	Method	Predicted beginning nucleotide location corresponding to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion
3259	A .	3	964	QMEPGNDTQISEFLLLGFSQEPGLQPFLFGLFLSM YLVTVLGNLLIILATISDSHLHTPMYFFLSNLSFA DICVTSTTIPKMLMNIQTQNKVITYIACLMQMYF FILFAGFENFLLSVMAYDRFVAICHPLHYMVIMN PHLCGLLVLASWTMSALYSLLQILMVVRLSFCT ALEIPHFFCELNQVIQLACSDSFLNHMVIYFTVAL LGGGPLTGILYSYSKIISSIHAISSAQGKYKAFSTC ASHLSVVSLFYGAILGVYLSSAATRNSHSSATAS VMYTVVTPMLNPFIYSLRNKDIKRALGIHLLWGT MKGQFFKKCP
3260	A	34	2573	IPFLKSCCCCCLFDFPPPPLDQVQEEECEVERVTE HGTPKPFRKFDSVAFGESQSEDEQFENDLETDPP NWQQLVSREVLLGLKPCEIKRQEVINELFYTERA HVRTLKVLDQVFYQRVSREGILSPSELRKIFSNLE DILQLHIGLNEQMKAVRKRNETSVIDQIGEDLLT WFSGPGEEKLKHAAATFCSNQPFALEMIKSRQK KDSRFQTFVQDAESNPLCRRLQLKDIIPTQMQRL TKYPLLLDNIATYTEWPTEREKVKKAADHCRQIL NYVNQAVKEAENKQRLEDYQRRLDTSSLKLSEY PNVEELRNLDLTKRKMIHEGPLVWKVNRDKTID LYTLLLEDILVLLQKQDDRLVLRCHSKILASTAD SKHTFSPVIKLSTVLVRQVATDNKALFVISMSDN GAQIYELVAQTVSEKTVWQDLICRMAASVKEQS TKPIPLPQSTPGEGDNDEEDPSKLKEEQHGISVTG LQSPDRDLGLESTLISSKPQSHSLSTSGKSEVRDL FVAERQFAKEQHTDGTLKEVGEDYQIAIPDSHLP VSEERWALDALRNLGLLKQLLVQQLGLTEKSVQ EDWQHFPRYRTASQGPQTDSVIQNSENIKAYHSG EGHMPFRTGTGDIATCYSPRTSTESFAPRDSVGL APQDSQASNILVMDHMIMTPEMPTMEPEGGLDD SGEHFFDAREAHSDENPSEGDGAVNKEEKDVNL RISGNYLILDGYDPVQESSTDEEVASSLTLQPMT GIPAVESTHQQQHSPQNTHSDGAISPFTPEFLVQQ RWGAMEYSCFEIQSPSSCADSQSQIMEYIHKIEA DLEHLKKVEESYTILCQRLAGSALTDKHSDKS
3261	A		2100	AVEFAEGALTMAPWPELGDAQPNPDKYLEGAA GQQPTAPDKSKETNKTDNTEAPVTKIELLPSYST ATLIDEPTEVDDPWNLPTLQDSGIKWSERDTKGK ILCFFQGIGRLILLLGFLYFFVCSLDILSSAFQLVG GKMAGQFFSNSSIMSNPLLGLVIGVLVTVLVQSS STSTSIVVSMVSSSLLTVRAAIPIIMGANIGTSITNT IVALMQVGDRSEFRRAFAGATVHDFFNWLSVLV LLPVEVATHYLEIITQLIVESFHFKNGEDAPDLLK VITKPFTKLIVQLDKKVISQIAMNDEKAKNKSLV KIWCKTFTNKTQINVTVPSTANCTSPSLCWTDGI QNWTMKNVTYKENIAKCQHIFVNFHLPDLAVGT ILLILSLLVLCGCLIMIVKILGSVLKGQVATVIKKT INTDFPFFFAWLTGYLAILVGAGMTFIVQSSSVFT SALTPLIGIGVITIERAYPLTLGSNIGTTTTAILAAL ASPGNALRSSLQIALCHFFFNISGILLWYPIPFTRL PIRMAKGLGNISAKYRWFAVFYLIIFFFLIPLTVFG LSLAGWRVLVGVGVPVVFIIILVLCLRLLQSRCPR VLPKKLQNWNFLPLWMRSLKPWDAVVSKFTGC FQMRCCCCCRVCCRACCLLCGCPKCCRCSKCCE DLEEAQEGQDVPVKAPETFDNITISREAQGEVPA

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			1055	SDSKTECTAL
3262	A		1377	SQQGSQPHRQGPPSLLTAPHSLDLPALPPGPRGS QGKLRRVLVPMSVKPSWGPGPSEGVTAVPTSDL GEIHNWTELLDLFNHTLSECHVELSQSTKRVVLF ALYLAMFVVGLVENLLVICVNWRGSGRAGLMN LYILNMAIADLGIVLSLPVWMLEVTLDYTWLWG SFSCRFTHYFYFVNMYSSIFFLVCLSVDRYVTLTS ASPSWQRYQHRVRRAMCAGIWVLSAIIPLPEVV HIQLVEGPEPMCLFMAPFETYSTWALAVALSTTI LGFLLPFPLITVFNVLTACRLRQPGQPKSRRHCLL LCAYVAVFVMCWLPYHVTLLLLTLHGTHISLHC HLVHLLYFFYDVIDCFSMLHCVINPILYNFLSPHF RGRLLNAVVHYLPKDQTKAGTCASSSSCSTQHSI IITKGDSQPAAAAPHPEPSLSFQAHHLLPNTSPISP TQPLTPS
3263	A	1	919	QARSPSVAAMASPQLCRALVSAQWVAEALRAP RAGQPLQLLDASWYLPKLGRDARREFEERHIPG AAFFDIDQCSDRTSPYDHMLPGAEHFAEYAGRL GVGAATHVVIYDASDQGLYSAPRVWWMFRAFG HHAVSLLDGGLRHWLRQNLPLSSGKSQPAPAEF RAQLDPAFIKTYEDIKENLESRRFQVVDSRATGR FRGTEPEPRDGIEPGHIPGTVNIPFTDFLSQEGLEK SPEEIRHLFQEKKVDLSKPLVATCGSGVTACHVA LGAYLCGKPDVPIYDGSWVEWYMRARPEDVISE GRGKTH
3264	A	1	1398	ARRSTPRTAPRASATRSAAGTMREIVHIQAGQCG NQIGAKFWEVISDEHGIDPTGSYHGDSDLQLERI NVYYNEAAGNKYVPRAILVDLEPGTMDSVRSGP FGQIFRPDNFVFGQSGAGNNWAKGHYTEGAELV DSVLDVVRKESESCDCLQGFQLTHSLGGGTGSG MGTLLISKIREEYPDRIMNTFSVMPSPKVSDTVVE PYNATLSVHQLVENTDETYSIDNEALYDICFRTL KLTTPTYGDLNHLVSATMSGVTTCLRFPGQLNA DLRKLAVNMVPFPRLHFFMPGFAPLTSRGSQQY RALTVPELTQQMFDSKNMMAACDPRHGRYLTV AAIFRGRMSMKEVDEQMLNVQNKNSSYFVEWIP NNVKTAVCDIPPRGLKMSATFIGNSTAIQELFKRI SEQFTAMFRRKAFLHWYTGEGMDEMEFTEAES NMNDLVSEYQQYQDATADEQGEFEEEEGEDEA
3265	A	265	862	WWEDARVLGPFHPEEEGHWVMTPSEGARAGTG RELEMLDSLLALGGLVLLRDSVEWEGRSLLKAL VKKSALCGEQVHILGCEVSEEEFREGFDSDINNR LVYHDFFRDPLNWSKTEEAFPGGPLGALRAMCK RTDPVPVTIALDSLSWLLLRLPCTTLCQVLHAVS HQDSCPGETPPSLFPLIHLPLPRSVPLFLSTLE
3266	A	802	1011	AAGAGADGREPASERASRAEPPAVAMGQNDLM GTAEDFADQFLRVTKQYLPHVARLCLISTFLEDG IRMWFQWSEQRDYIDTTWNCGYLLASSFVFLNL LGQLTGCVLVLSRNFVQYACFGLFGIIALQTIAYS ILWDLKFLMRNLALGGGLLLLLAESRSEGKSMF AGVPTMRESSPKQYMQLGGRVLLVLMFMTLLH FDASFFSIVQNIVGTALMILVAIGFKTKLAALTLV VWLFAINVYFNAFWTIPVYKPMHDFLKYDFFQT MSVIGGLLLVVALGPGGVSMDEKKKEW ASTFCSAWKRRSTAALWWSGSRASRSHPRELGP

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				LCFVFGTAALSIRSMDVLSLFLEHGKLVFASGLSP RA
3268	A	490	679	EDAWITNPSLSNARSTPSKPLCYTVLKEGQVVGV KTTKASNTREKLRPESERRMVKSFGDEVT
3269	A	2	796	GSTHASGARPSLKRARSQRGRPLPSRALPSAHKD MTTNAGPLHPYWPQHLRLDNFVPNDRPTWHILA GLFSVTGVLVVTTWLLSGRAAVVPLGTWRRLSL CWFAVCGFIHLVIEGWFVLYYEDLLGDQAFLSQ LWKEYAKGDSRYILGDNFTVCMETITACLWGPL SLWVVIAFLRQHPLRFILQLVVSVGQIYGDVLYF LTEHRDGFQHGELGHPLYFWFYFVFMNALWLV LPGVLVLDAVKHLTHAQSTLDAKATKAKSKKN
3270	A	17	229	GDTGPQILMSYLDSVASKLLQMVKKLSQSFCSNF KYLTKYSRKQVSDEIKKSRRTVESNPIFFKKNKKI Q
3271	A	419	553	IQSGLSLCFADLSETPEGRAGVPGCPHSCDGVAS GRPCSPSSAG
3272	A	1211	1450	FQFIQIELLNILQSLIRNQTQSPYNTTAYPAIDSVIT ILPFSFSCFFIITKCFGLSIFPSVIFFLHVYFILTLVVF YCC
3273	A	59	1562	QAWSLQVALSPFFPASPSNSFAAAVPQLLFPELP LPHVPGQESAKRRSARRFLIMSELTKELMELVW GTKSSPGLSDTIFCRWTQGFVFSESEGSALEQFEG GPCAVIAPVQAFLLKKLLFSSEKSSWRDCSQEEQ KELLCHTLCDILESACCDHSGSYCLVSWLRGKTT EETASISGSPÄESSCQVEHSSALAVEELGFERFHA LIQKRSFRSLPELKDAVLDQYSMWGNKFGVLLF LYSVLLTKGIENIKNEIEDASEPLIDPVYGHGSQS LINLLLTGHAVSNVWDGDRECSGMKLLGIHEQA AVGFLTLMEALRYCKVGSYLKISKIPYLDCLASE THLTVFFAKDMALVAPEAPSEQARRVFQTYDPE DNGFIPDSLLEDVMKALDLVSDPEYINLMKNKL DPEGLGIILLGPFLQEFFPDQGSSGPESFTVYHYN GLKQSNYNEKVMYVEGTAVVMGFEDPMLQTD DTPIKRCLQTKWPYIELLWTTDRSPSLN
3274	A	186	1358	RVVHRFFKSSAFWPAEVKQPRGGPKTGSRKEGA GSRAPQPVVRSFCGSVGAEGRMEKLRLLGLRYQ EYVTRHPAATAQLETAVRGFSYLLAGRFADSHE LSELVYSASNLLVLLNDGILRKELRKKLPVSLSQ QKLLTWLSVLECVEVFMEMGAAKVWGEVGRW LVIALIQLAKAVLRMLLLLWFKAGLQTSPPIVPL DRETQAQPPDGDHSPGNHEQSYVGKRSNRVVRT LQNTPSLHSRHWGAPQQREGRQQQHHEELSATP TPLGLQETIAEFLYIARPLLHLLSLGLWGQRSWK PWLLAGVVDVTSLSLLSDRKGLTRRERRELRRR TILLLYYLLRSPFYDRFSEARILFLLQLLADHVPG VGLVTRPLMDYLPTWQKIYFYSWG
3275	A	575	759	SVYSASSCKCCNYRKTEQIPDCEQPPASSMPERPS HESQPTPQMMPLSAPSRAEELGQRPG
3276	A	7	258	KAAGHRLLLAAGHPSMPSSDCLLWEGSLELRPL QHISSLLVLVSTTCLFAFPRVPIAFESKSCLIYHCH CAFTVRHYMCSSHTG
3277	A	9	2221	KLGVEPEEEGGGDDEEDAEAWAMELADVGAAA SSQGVHDQVLPTPNASSRVIVHVDLDCFYAQVE MISNPELKDKPLGVQQKYLVVTCNYEARKLGVK

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		to first amino acid residue of peptide	acid residue of peptide sequence	X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion
ļ		sequence		KLMNVRDAKEKCPQLVLVNGEDLTRYREMSYK
		ĺ		VTELLEEFSPVVERLGFDENFVDLTEMVEKRLQQ
				LOSDELSAVTVSGHVYNNQSINLLDVLHIRLLVG
				SQIAAEMREAMYNQLGLTGCAGVASNKLLAKL
				VSGVFKPNQQTVLLPESCQHLIHSLNHIKEIPGIG
İ				YKTAKCLEALGINSVRDLQTFSPKILEKELGISVA
				QRIQKLSFGEDNSPVILSGPPQSFSEEDSFKKCSSE
				VEAKNKIEELLASLLNRLCQDERKPHTVRLIIRRY
				SSEKHYGRESRQCPIPSHVIQKLGTGNYDVMTPM
				VDILMKLFRNMVNVKMPFHLTLLSVCFCNLKAL
1				NTAKKGLIDYYLMPSLSTTSRSGKHSFKMKDTH
		1		MEDFPKDKETNRDFLPSGRIESTRTRESPLDTTNF
,				SKEKDINEFPLCSLPEGVDQEVFKQLPVDIQEEIL
		[SGKSREKFQGKGSVSCPLHASRGVLSFFSKKQM
				QDIPINPRDHLSSSKQVSSVSPCEPGTSGFNSSSSS YMSSQKDYSYYLDNRLKDERISQGPKEPQGFHF
}			ļ	TNSNPAVSAFHSFPNLQSEQLFSRNHTTDSHKQT
				VATDSHEGLTENREPDSVDEKITFPSDIDPQVFYE
	Ĭ	1		LPEAVQKELLAEWKRTGSDFHIGHK
3278	A	1	876	GLRLHVDLVEKPRTGIMAAETRNVAGAEAPPPQ
32.0]	•	.	KRYYRORAHSNPMADHTLRYPVKPEEMDWSEL
				YPEFFAPLTQNQSHDDPKDKKEKRAQAQVEFAD
	,		-	IGCGYGGLLVELSPLFPDTLILGLEIRVKVSDYVQ
,			ļ	DRIRALRAAPAGGFQNIACLRSNAMKHLPNFFY
ĺ				KGQLTKMFFLFPDPHFKRTKHKWRIISPTLLAEY
				AYVLRVGGLVYTITDVLELHDWMCTHFEEHPLF
			•	ERVPLEDLSEDPVVGHLGTSTEEGKKVLRNGGK
2070		90	2020	NFPAIFRRIQDPVLQAVTSQTSLPGH TRTKRRLGREKAMASPPRGWGCGELLLPFMLLG
3279	A	82	2929	TLCEPGSGQIRYSMPEELDKGSFVGNIAKDLGLE
		}		PQELAERGVRIVSRGRTQLFALNPRSGSLVTAGRI
				DREELCAQSPLCVVNFNILVENKMKIYGVEVEII
				DINDNFPRFRDEELKVKVNENAAAGTRLVLPFA
				RDADVGVNSLRSYQLSSNLHFSLDVVSGTDGQK
[ĺ	ĺ	YPELVLEQPLDREKETVHDLLLTALDGGDPVLSG
				TTHIRVTVLDANDNAPLFTPSEYSVSVPENIPVGT
'				RLLMLTATDPDEGINGKLTYSFRNEEEKISETFQL
				DSNLGEISTLQSLDYEESRFYLMEVVAQDGGAL
	1			VASAKVVVTVQDVNDNAPEVILTSLTSSISEDCL
				PGTVIALFSVHDGDSGENGEIACSIPRNLPFKLEK SVDNYYHLLTTRDLDREETSDYNITLTVMDHGT
			1	PPLSTESHIPLKVADVNDNPPNFPQASYSTSVTEN
	1			NPRGVSIFSVTAHDPDSGDNARVTYSLAEDTFQG
]	1		APLSSYVSINSDTGVLYALRSFDYEQLRDLQLWV
				TASDSGNPPLSSNVSLSLFVLDQNDNTPEILYPAL
				PTDGSTGVELAPRSAEPGYLVTKVVAVDKDSGQ
	ļ			NAWLSYRLLKASEPGLFAVGLHTGEVRTARALL
]			DRDALKQSLVVAVEDHGQPPLSATFTVTVAVAD
				RIPDILADLGSIKTPIDPEDLDLTLYLVVAVAAVS
				CVFLAFVIVLLVLRLRRWHKSRLLQAEGSRLAG
	1			VPASHFVGVDGVRAFLQTYSHEVSLTADSRKSH
				LIFPQPNYADTLLSEESCEKSEPLLMSDKVDANK
		Į	}	EERRVQQAPPNTDWRFSQAQRPGTSGSQNGDDT
				GTWPNNQFDTEMLQAMILASASEAADGSSTLGG
L	<u> </u>	L	L	GAGTMGLSARYGPQFTLQHVLQGELGSDYRQN

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				VYIPGSNATLTNAAGKRDGKAPAGGNGNKKKS GKKEKK
3280	A	149		GTSQMSSHKGSVVAQGNGAPASNREADTAELAE LGPLLEEKGKRVIANPPKAEEEQTCPVPQEEEEE VRVLTLPLQAHHAMEKMEEFVYKVWEGRWRVI PYDVLPDWLKDNDYLLHGHRPPMPSFRACFKSIF RIHTETGNIWTHLLGFVLFLFLGILTMLRPNMYF MAPLQEKVVFGMFFLGAVLCLSFSWLFHTVYCH SEKVSRTFSKLDYSGIALLIMGSFVPWLYYSFYCS PQPRLIYLSIVCVLGISAIIVAQWDRFATPKHRQT RAGVFLGLGLSGVVPTMHFTIAEGFVKATTVGQ MGWFFLMAVMYITGAGLYAARIPERFFPGKFDI WFQSHQIFHVLVVAAAFVHFYGVSNLQEFRYGL EGGCTDDTLL
3281	A	1	557	RPRRRQPSFSCRVLVLEDPPCFRFTNSMNQEKLA KLQAQVRIGGKGTARRKKKVVHRTATADDKKL QSSLKKLAVNNIAGIEEVNMIKDDGTVIHFNNPK VQASLSANTFAITGHAEAKPITEMLPGILSQLGAD SLTSLRKLAEQFPRQVLDSKAPKPEDIDEEDDDV PDLVENFDEASKNEAN
3282	A	155	1139	HALGRRGGSQELSAAACGCFALRLRAPGSGRPA LAPGAAAFAGLGGAPRFPPRGSAAGRTMLLKEY RICMPLTVDEYKIGQLYMISKHSHEQSDRGEGVE VVQNEPFEDPHHGNGQFTEKRVYLNSKLPSWAR AVVPKIFYVTEKAWNYYPYTITEYTCSFLPKFSIH IETKYEDNKGSNDTIFDNEAKDVEREVCFIDIACD EIPERYYKESEDPKHFKSEKTGRGQLREGWRDSH QPIMCSYKLVTVKFEVWGLQTRVEQFVHKVVR DILLIGHRQAFAWVDEWYDMTMDDVREYEKN MHEQTNIKVCNQHSSPVDDIESHAQTST
3283	A	159	547	IKSKLNQQVEVQESEWRLTEAKGPTMGKESGW DSGRAAVAAVVGGVVAVGTVLVALSAMGFTSV GIAASSIAAKMMSTAAIANGGGVAAGSLVAILQS VGAAGLSVTSKVIGGFAGTALGAWLGSPPSS
3284	A	227	637	TSNSLLRPDRMSVMDLANTCSSFQSDLDFCSDCG SVLPLPGAQDTVTCIRCGFNINVRDFEGKVVKTS VVFHQLGTAMPMSVEEGPECQGPVVDRRCPRCG HEGMAYHTRQMRSADEGQTVFYTCTNCKFQEK EDS
3285	A	123	1535	HRLSYDEAFAMANDPLEGFHEVNLASPTSPDLL GVYESGTQEQTTSPSVIYRPHPSALSSVPIQANAL DVSELPTQPVYSSPRRLNCAEISSISFHVTDPAPCS TSGVTAGLTKLTTRKDNYNAEREFLQGATITEAC DGSDDIFGLSTDSLSRLRSPSVLEVREKGYERLKE ELAKAQRELKLKDEECERLSKVRDQLGQELEEL TASLFEEAHKMVREANIKQATAEKQLKEAQGKI DVLQAEVAALKTLVLSSSPTSPTQEPLPGGKTPF KKGHTRNKSTSSAMSGSHQDLSVIQPIVKDCKEA DLSLYNEFRLWKDEPTMDRTCPFLDKIYQEDIFP CLTFSKSELASAVLEAVENNTLSIEPVGLQPIRFV KASAVECGGPKKCALTGQSKSCKHRIKLGDSSN YYYISPFCRYRITSVCNFFTYIRYIQQGLVKQQDV DQMFWEVMQLRKEMSLAKLGYFKEEL
3286	Α	3	589	GPSQSMAAGELEGGKPLSGLLNALAQDTFHGYP GITEELLRSQLYPEVPPEEFRPFLAKMRGILKSIAS

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				ADMDFNQLEAFLTAQTKKQGGITSDQAAVISKF WKSHKTKIRESLMNQSRWNSGLRGLSWRVDGK SQSRHSAQIHTPVAIIELELGKYGQESEFLCLEFD EVKVNQILKTLSEVEESISTLISQPN
3287	A	50	390	LGAMAKHHPDLIFCRKQAGVAIGRLCEKCDGKC VICDSYVRPCTLVRICDECNYGSYQGRCVICGGP GVSDAYYCKECTIQEKDRDGCPKIVNLGSSKTDL FYERKKYGFKKR
3288	A	3	428	RTTFFRFRPCESLCGDMKLLTHNLLSSHVRGVGS RGFPLRLQATEVRICPVEFNPNFVARMIPKVEWS AFLEAADNLRLIQVPKGPVEGYEENEEFLRTMH HLLLEVEVIEGTLQCPESGRMFPISRGIPNMLLSE EETES
3289	A		1743	AGCCRDTRFPTPRGPGSLCHNFCRSAACTVTRTI HGSPREDTGTPRSREMMFQDSVAFEDVAVSFTQ EEWALLDPSQKNLYRDVMQETFKNLTSVGKTW KVQNIEDEYKNPRRNLSLMREKLCESKESHHCG ESFNQIADDMLNRKTLPGITPCESSVCGEVGTGH SSLNTHIRADTGHKSSEYQEYGENPYRNKECKK AFSYLDSFQSHDKACTKEKPYDGKECTETFISHS CIQRHRVMHSGDGPYKCKFCGKAFYFLNLCLIH ERIHTGVKPYKCKQCGKAFTRSTTLPVHERTHTG VNADECKECGNAFSFPSEIRRHKRSHTGEKPYEC KQCGKVFISFSSIQYHKMTHTGEKPYECKQCGK AFRCGSHLQKHGRTHTGEKPYECRQCGKAFRCT SDLQRHEKTHTEDKPYGCKQCGKGFRCASQLQI HERTHSGEKPHECKECGKVFKYFSSLRIHERTHT GEKPHECKQCGKAFRYFSSLHIHERTHTGDKPYE CKVCGKAFTCSSSIRYHERTHTGEKPYECKHCGK AFISNYIRYHERTHTGEKPYQCKQCGKAFIRASS
3290	A		1350	GRPRSSSDNRNFLRERAGLSSAAVQTRIGNSAAS RRSPAARPPVPAPPALPRGRPGTEGSTSLSAPAVL VVAVAVVVVVSAVAWAMANYIHVPPGSPEVP KLNVTVQDQEEHRCREGALSLLQHLRPHWDPQE VTLQLFTDGITNKLIGCYVGNTMEDVVLVRIYGN KTELLVDRDEEVKSFRVLQAHGCAPQLYCTFNN GLCYEFIQGEALDPKHVCNPAIFRLIARQLAKIHA IHAHNGWIPKSNLWLKMGKYFSLIPTGFADEDIN KRFLSDIPSSQILQEEMTWMKEILSNLGSPVVLCH NDLLCKNIIYNEKQGDVQFIDYEYSGYNYLAYDI GNHFNEFAGVSDVDYSLYPDRELQSQWLRAYLE AYKEFKGFGTEVTEKEVEILFIQVNQFALASHFF WGLWALIQAKYSTIEFDFLGYAIVRFNQYFKMK PEVTALKVPE
3291	A	102	839	PEAQTSAVLAREKGHLPTMRHEAPMQMASAQD ARYGQKDSSDQNFDYMFKLLIIGNSSVGKTSFLF RYADDSFTSAFVSTVGIDFKVKTVFKNEKRIKLQI WDTAGQERYRTITTAYYRGAMGFILMYDITNEE SFNAVQDWSTQIKTYSWDNAQVILVGNKCDME DERVISTERGQHLGEQLGFEFFETSAKDNINVKQ TFERLVDIICDKMSESLETDPAITAAKQNTRLKET PPPPQPNCAC
3292	A	2	4136	DRPPWNSRVDDFVTNLIHLSSKGHISPAKDTSLQ QRTPAEMSPVLHFYVRPSGHEGAASGHTRRKLQ

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				GKLPELQGVETELCYNVNWTAEALPSAEETKKL MWLFGCPLLLDDVARESWLLPGSNDLLLEVGPR LNFSTPTSTNIVSVCRATGLGPVDRVETTRRYRLS FAHPPSAEVEAIALATLHDRMTEQHFPHPIQSFSP ESMPEPLNGPINILGEGRLALEKANQELGLALDS WDLDFYTKRFQELQRNPSTVEAFDLAQSNSEHS RHWFFKGQLHVDGQKLVHSLFESIMSTQESSNP NNVLKFCDNSSAIQGKEVRFLRPEDPTRPSRFQQ QQGLRHVVFTAETHNFPTGVCPFSGATTGTGGRI RDVQCTGRGAHVVAGTAGYCFGNLHIPGYNLP WEDLSFQYPGNFARPLEVAIEASNGASDYGNKF GEPVLAGFARSLGLQLPDGQRREWIKPIMFSGGI GSMEADHISKEAPEPGMEVVKVGGPVYRIGVGG GAASSVQVQGDNTSDLDFGAVQRGDPEMEQKM NRVIRACVEAPKGNPICSLHDQGAGGNGNVLKE LSDPAGAIIYTSRFQLGDPTLNALEIWGAEYQESN ALLLRSPNRDFLTHVSARERCPACFVGTITGDRRI VLVDDRECPVRRNGQGDAPPTPPPTPVDLELEW VLGKMPRKEFFLQRKPPMLQPLALPPGLSVHQA LERVLRLPAVASKRYLTNKVDRSVGGLVAQQQC VGPLQTPLADVAVVALSHEELIGAATALGEQPV KSLLDPKVAARLAVAEALTNLVFALVTDLRDVK CSGNWMWAAKLPGEGAALADACEAMVAVMA ALGVAVDGGKDSLSMAARVGTETVRAPGSLVIS AYAVCPDITATVTPDLKHPEGRGHLLYVALSPG QHRLGGTALAQCFSQLGEHPPDLDLPENLVRAFS ITQGLLKDRLLCSGHDVSDGGLVTCLLEMAFAG NCGLQVDVPVPRVDVLSVLFAEEPGLVLEVQEP DLAQVLKRYRDAGLHCLELGHTGEAGPHAMVR VSVNGAVVLEEPVGELRALWEETSFQLDRLQAE PRCVAEEERGLRERMGPSYCLPPTFPKASVPREP GGPSPRVAILREEGSNGDREMADAFHLAGFEVW DVTMQDLCSGAIGLDTFRGVAFVGGFSYADVLG SAKGWAAAVTFHPRAGAELRRFRKRPDTFSLGV CNGCQLLALLGWVGGDPNEDAAEMGPDSQPAR PGLLLRHNLSGRYESRWASVRVGPGPALMLRG MEGAVLPVWSAHGEGYVAFSSPELQAQIEARGL APLHWADDDGNPTEQYPLNPNGSPGGVAGICSC DGRHLAVMPHPERAVRPWQWAWRPPPFDTLTT SPWLQLFINARNWTLEGSC
3293	A	65	642	GVRGFWAGTMASRAGPRAAGTDGSDFQHRERV AMHYQMSVTLKYEIKKLIYVHLVIWLLLVAKMS VGHLRLLSHDQVAMPYQWEYPYLLSILPSLLGLL SFPRNNISYLVLSMISMGLFSIAPLIYGSMEMFPA AQQLYRHGKAYRFLFGFSAVSIMYLVLVLAVQV HAWQLYYSKKLLDSWFTSTQEKKHK
3294	A	35	1821	SQRSCPRSPSSPAPPWARCSNPDSRTGGVPVPRA WSAGGPALGLMAAPVRLGRKRPLPACPNPLFVR WLTEWRDEATRSRHRTRFVFQKALRSLRRYPLP LRSGKEAKILQHFGDGLCRMLDERLQRHRTSGG DHAPDSPSGENSPAPQGRLAEVQDSSMPVPAQP KAGGSGSYWPARHSGARVILLVLYREHLNPNGH HFLTKEELLQRCAQKSPRVAPGSARPWPALRSLL HRNLVLRTHQPARYSLTPEGLELAQKLAESEGLS LLNVGIGPKEPPGEETAVPGAASAELASEAGVQQ

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	·			QPLELRPGEYRVLLCVDIGETRGGGHRPELLREL QRLHVTHTVRKLHVGDFVWVAQETNPRDPANP GELVLDHIVERKRLDDLCSSIIDGRFREQKFRLKR CGLERRVYLVEEHGSVHNLSLPESTLLQAVTNTQ VIDGFFVKRTADIKESAAYLALLTRGLQRLYQGH TLRSRPWGTPGNPESGAMTSPNPLCSLLTFSDFN AGAIKNKAQSVREVFARQLMQVRGVSGEKAAA LVDRYSTPASLLAAYDACATPKEQETLLSTIKCG RLQRNLGPALSRTLSQLYCSYGPLT
3295	A	2	1115	EFHPHTQVSGLTTPQLQEPDVWSPSRGQPVSLHL PGKGAPEVKEMAWWKSWIEQEGVTVKSSSHFN PDPDAETLYKAMKGIGTNEQAIIDVLTKRSNTQR QQIAKSFKAQFGKDLTETLKSELSGKFERLIVAL MYPPYRYEAKELHDAMKGLGTKEGVIIEILASRT KNQLREIMKAYEEDYGSSLEEDIQADTSGYLERI LVCLLQGSRDDVSSFVDPALALQDAQDLYAAGE KIRGTDEMKFITILCTRSATHLLRVFEEYEKIANK SIEDSIKSETHGSLEEAMLTVVKCTQNLHSYFAE RLYYAMKGAGTRDGTLIRNIVSRSEIDLNLIKCH FKKMYGKTLSSMIMEDTSGDYKNALLSLVGSDP
3296	A	1	838	GTRGGVGPGDNGGVEAGAKPGAAAIPLRGDGS GETGPGRVAPGEVRGSPRGHVAGPEGPREVLFFF FLPSSKPASEVINEYSWKVDFLKGMLQAEKLTSS SEKALANQFLAPGRVPTTARERVPATKTVHLQS RARYTSEMRSELLGTDSAEPEMDVRKRTGVAGS QPVSEKQSAAELDLVLQRHQNLQEKLAEEMLGL ARSLKTNTLAAQSVIKKDNQTLSHSLKMADQNL EKLKTESERLEQHTQKSVNWLLWAMLIIVCFIFIS MILFIRIMPKLK
3297	A	46	617	HKQPAGFLGLWLGTETYTISFPGPETFGLGLSHA TGIPGSPACRQPVVGLHSLHNYRMAMVSAMSW VLYLWISACAMLLCHGSLQHTFQQHHLHRPEGG TCEVIAAHRCCNKNRIEERSQTVKCSCLPGKVAG TTRNRPSCVDASIVIGKWWCEMEPCLEGEECKTL PDNSGWMCATGNKIKTTRIHPRT
3298	A	157	748	IQPPDPRNMTLAAYKEKMKELPLVSLFCSCFLAD PLNKSSYKYEADTVDLNWCVISDMEVIELNKCT SGQSFEVILKPPSFDGVPEFNASLPRRRDPSLEEIQ KKLEAAEERRKYQEAELLKHLAEKREHEREVIQ KAIEENNNFIKMAKEKLAQKMESNKENREAHLA AMLERLQEKDKHAEEVRKNKELKEEASR
3299	A			TQLPAPLSGVLSRLQLGSGAPLLTWVQETAGVA GGAPRRTPVTMWRLLARASAPLLRVPLSDSWA LLPASAGVKTLLPVPSFEDVSIPEKPKLRFIERAPL VPKVRREPKNLSDIRGPSTEATEFTEGNFAILALG GGYLHWGHFEMMRLTINRSMDPKNMFAIWRVP APFKPITRKSVGHRMGGGKGAIDHYVTPVKAGR LVVEMGGRCEFEEVQGFLDQVAHKLPFAAKAVS RGTLEKMRKDQEERERNNQNPWTFERIATANML GIRKVLSPYDLTHKGKYWGKFYMPKRV
3300	A	2	1847	FVAGGPRGSGSAAETMPEIRVTPLGAGQDVGRS CILVSIAGKNVMLDCGMHMGFNDDRRFPDFSYI TQNGRLTDFLDCVIISHFHLDHCGALPYFSEMVG YDGPIYMTHPTQAICPILLEDYRKIAVDKKGEAN FFTSQMIKDCMKKVVAVHLHQTVQVDDELEIKA

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				YYAGHVLGAAMFQIKVGSESVVYTGDYNMTPD RHLGAAWIDKCRPNLLITESTYATTIRDSKRCRE RDFLKKVHETVERGGKVLIPVFALGRAQELCILL ETFWERMNLKVPIYFSTGLTEKANHYYKLFIPWT NQKIRKTFVQRNMFEFKHIKAFDRAFADNPGPM VVFATPGMLHAGQSLQIFRKWAGNEKNMVIMP GYCVQGTVGHKILSGQRKLEMEGRQVLEVKMQ VEYMSFSAHADAKGIMQLVGQAEPESVLLVHGE AKKMEFLKQKIEQELRVNCYMPANGETVTLPTS PSIPVGISLGLLKREMAQGLLPEAKKPRLLHGTLI MKDSNFRLVSSEQALKELGLAEHQLRFTCRVHL HDTRKEQETALRVYSHLKSVLKDHCVQHLPDGS VTVESVLLQAAAPSEDPGTKVLLVSWTYQDEEL GSFLTSLLKKGLPQAPS
3301	A	2	349	CIRTEPAAAFRRLGALSGAAALGFASYGAHGAQ FPDAYGKELFDKANKHHFLHSLALLGVPHCRKP LWAGLLLASGTTLFCTSFYYQALSGDPSIQTLAP AGGTLLLLGWLALAL
3302	A	59	1184	LRRNCSALGGLFQTIISDMKGSYPVWEDFINKAG KLQSQLRTTVVAAAAFLDAFQKVADMATNTRG GTREIGSALTRMCMRHRSIEAKLRQFSSALIDCLI NPLQEQMEEWKKVANQLDKDHAKEYKKARQEI KKKSSDTLKLQKKAKKGRGDIQPQLDSALQDVN DKYLLLEETEKQAVRKALIEERGRFCTFISMLRP VIEEEISMLGEITHLQTISEDLKSLTMDPHKLPSSS EQVILDLKGSDYSWSYQTPPSSPSTTMSRKSSVC SSLNSVNSSDSRSSGSHSHSPSSHYRYRSSNLAQQ APVRLSSVSSHDSGFISQDAFQSKSPSPMPPEAPN QRRKEKREPDPNGGGPTTASGPPAAAEEAQRPRS M
3303	A :	511	958	AGRGGPGKPVSWSSGPGSPGQTQRRSWVKSTRG HSSLLPPSQDFVAGLSVILRGTVDDRLNWAFNLY DLNKDGCITKEEMLDIMKSIYDMMGKYTYPALR EEAPREHVESFFQKMDRNKDGVVTIEEFIESCQK DENIMRSMQLFDNVI
3304	A	40	432	ISEAASGAFQAR*FYQMLEQKTDALGKQSVNRG FTKDKTLSSIFNIEMVKEKTAEEIKQIWQQYFAA KDTVYAVIPAEKFDLIWNRAQSCPTFLCALPRRE GYEFFVGQWTGTELHFHCTYKYSDPEGKA
3305	A	2	483	LDACSTGPYSRSTHASADAWADAWVVVVLKVV GMTLFLLYFPQIFNKSNDGFTTTRSYGTVSQIFGS RSPSPNGFITTRSYGTVCPKDWEFYQARCFFLIHL *\SSWNESWDFCKGKGCTLAIVDNSETLKLLHDL HDAEKNYIALPYRSSKYMSTCNGTF
3306	A	2	872	TLSSACLIGDAWKELTIVAGAVSNQLLVWYPAT ALADNKPVAPDRRISGHVGIIFSMSYLESKGLLA TASEDRSVRIWKGGDLRVPGGRVQNIGHCFGHS ARVWQVKLLENYLISAGEDCVCLVWSHEGEILQ AFRGHQGRGIRAIAAHERQAWVITGGDDSGIRL WHLVGRGYRGLG/DLGSLLQVP**ARYTQGCDS GWLLATAGSD*YRGPVSL*RRGQVLGAAARG*T FPVLLPAGGSSWSRGLRIVCYGQWGRSCQGCPH QHSNCCCGPDPVSWEGAQLELGPAWL
3307	A	2	927	RTSRVEKGLRKAGAAVTMESDEWFSQALPANTS AQKAELIALTQAIRWGKDINVNTDSRYAFATVH

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| Predicted beginning nucleotide location | Predicted end nucleotide | Predicted end nucleotide location | Predicted end nucleotide | Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine,

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			·	VRGAICQERRLLTSAEKAIKNKNPPSSKPNRSSS\F WGTTCDQVNAKQGPKPSPGHRLRRNLPGEKWEI DFTKVKPHQAGYKYLLVLVDTFSGWTEAFATK NETVNMVVKFLLNEIIPRHGLPVAIGSDNGPAFA LSIV*SVSKALNIQWKLHCAYRPQSSGQVERMNC TLKNTLTKLILETGVNWVSLLPLALLRVRCTPYW AGFLPFEIMYGRVLPILPKLRDAQLAKISQTNLLQ YLQSP
3308	A	490	1077	NSPSLDFNDNEDIPTELSDSSDTHDEGEVQAFYE DLSGRQYVNEVFNFSVDKLYDLLFTNSPFQRDF MEQRRFSDIIFHPWKKEENGNQSRVIPYTITLTNP LEHKTATVRETQTMYKASQESECYVIDAEVLTH DVPYHDYFYTINRYTLTRVARNKSRLRVSTELRY RKQPWGLVKTFIEKNFWSGLEDYFRHL
3309	A	490	1077	NSPSLDFNDNEDIPTELSDSSDTHDEGEVQAFYE DLSGRQYVNEVFNFSVDKLYDLLFTNSPFQRDF MEQRRFSDIIFHPWKKEENGNQSRVIPYTITLTNP LEHKTATVRETQTMYKASQESECYVIDAEVLTH DVPYHDYFYTINRYTLTRVARNKSRLRVSTELRY RKQPWGLVKTFIEKNFWSGLEDYFRHL
3310	A	2	1198	SPLCHPGLSRER/S*SEAKLRSGRYC*KRQVEAPL *RPGL*TMAASDTERDGLAPEKTSPDRDKKKEQS EVSVSPRASKHHYSRSRSRSRERKRKSDNEGRKH RSRSRSKEGRRHESKDKSSKKHKSEEHNDKEHSS DKGRERLNSSENGEDRHKRKERKSSRGRSHSRS RSRERRHRSRSRERKKSRSRSRERKKSRSRSRER
i				KKSRSRSRERKRRIRSRSRSRSRHRHRTRSRSRTR SRSRDRKKRIEKPRRFSRSLSRTPSPPPFRGRNTA MDAQEALARRLERAKKLQEQREKEMVEKQKQQ EIAAAAAATGGSVLNVAALLASGTQVTPQIAMA AQMAALQAKALAETGIAVPSYYNPAAVNPMKF AEQEKKRKMLWQGKKEGDKSQSAGNMGKN
3311	A	177	4	PIQIPPRITPPRPSPHLLTPRTGSSPPPPRAPSPPHPT PGPAHDFPPLSAVLSGHTKT
3312	A .	3	426	LESPRH*PPCWGPLIWALTVSSVPSPTPELSCILKS P/RPACPV/PGLWPSLLSPAPPQSSGPLLGLSPCPG AGQWPSPLSPAPPPSSDPLSGLSPCPGAGPRSSP\S ASAPCRAVPLSPRRLTWPPHLQVGILIPTGRPWK NL
3313	A	162	2	QLQNLASRGCL*SQLLRRLRRENRLNPGGGGCSE IAP\CTPAWVTQRDFFRKKK
3314	A	162	2	QLQNLASRGCL*SQLLRRLRRENRLNPGGGGCSE IAP\CTPAWVTQRDFFRKKK
3315	A	466	1	PRKRESWWGERLP/PRGFPPAAEDAPAPGWKGR KHASRTARAHVFHPIRQSIRSPVRGRPGDPRAAH TRSAGTRLQCKASRGG*GKGPAPTR*EGGPGSAP APLPASSGCSLFPDSSPWTPPPPAPGAAAAQP**T PRCPAALRAGAHIGRVGRPY
3316	A	3	2307	NHLGTLMQNWDSSSRVPFSSGQHSTQSFPPSLMS KSNSMLQKPT\AYVRPMDGQESMEPKLSSEHYSS QSHGNSMTELKPSSKAHLTKLKIPSQPLDASASG DVSCVDEILKEMTHSWPPPLTAIHTPCKTEPSKFP FPTKESQQSNFGTGEQKRYNPSKTSNGHQSKSM LKDDLKLSSSEDSDGEQDCDKTMPRSTPGSNSEP SHHNSEGADNSRDDSSSHSGSESSSGSDSESESSS

	1 52 2 2			
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	·			SDSEANEPSQSASPEPEPPPTNKWQLDNWLNKV NPHKVSPASSVDSNIPSSQGYKKEGREQGTGNSY TDTSGPKETSSATPGR\APKPIQKGSESGRGRQKS PAQSDSTTQRRTVGKKQPKKAEKAAAEEPRGGL KIESETPVDLASSMPSSRHKAATKGSRKPNIKKES KSSPRPTAEKKKYKSTSKSSQKSREIIETDTSSSDS DESESLPPSSQTPKYPESNRTPVKPSSVEEEDSFFR QRMFSPMEEKELLSPLSEPDDRYPLIVKIDLNLLT RIPGKPYKETEPPKGEKKNVPEKHTREAQKQASE KVSNKGKRKHKNEDDNRASESKKPKTEDKNSA GHKPSSNRESSKQSAAKEKDLLPSPAGPVPSKDP KTEHGSRKRTISQSSSLKSSSNSNKETSGSSKNSS STSKQKKTEGKTSSSSKEVKVKAPSSSSNCPPSAP TLDSSKPRRTKLVFDDRNYSADHYLQEAKKLKH NADALSDRFEKAVYYLDAVVSFIECGNALEKNA QESKSPFPMYSETVDLI
3317	A	496	2	NLLQDEKLVHSYPYDWRTQETCGYIVPARQWFI N\TRDIKTAAKELLKKVKFIPGSALNGMVEMMD RRPYWCISRQRVWGVPIPVFHHKTKDEYLINSQT TEHIVKLVEQHGSDIWWTLPPEQLLPKEVLSEVG GPDALEYVPGQDILDIWFDSGTSWSYVLPGPD
3318	A	2	512	AWHEGDSRSDQCHHPYNYGFDYYYGMPFTLVD SCWPDPSRNTELAFESQLWLCVQLVAIAILTLTF GKLSGWVSVPWLLIFSMILFIFLLGYAWFSSHTSP LYWDCLLMRGHEITEQPMKAE\RAGSIMVKEAIF LFRKGHSKGKLFLLFFLPFLQVHKTFPTTDGFHW AP
3319	A	407	1	SSLHRSPRPASPLPVPEAP\SFLPVPAPKPSALPPFS LSGAPSSASTFSPHSSPSPASPTPAPSPQSPFPSRPT SPPSLTPTRRPPLPADRRGPHLLYQPLHAPLEAAA TGPE/PSAAAGRLPRPRPPWRAAYPASR
3320	A	4037	3432	QMSEAVAEKMLQYRRDTAGWKICREGNGVSVS WRPSVEFPGNLYRGEGIVYGTLEEVWDCVKPAV GGLRVKWDENVTGFEIIQSITDTLCVSRTSTPSAA MKLISPRDFVDLVLVKRYEDGTISSNATHVEHPL CPPKPGFVRGFNHPCGCFCEPLPGEPTKTNLVTFF HTDLSGYLPQNVVDSFFPRSMTRFYANLQKAVK
3321	A	37	360	SHSASGAGRPAAPAADLRPAPNGQRPGPRLGAR ALWLPPRGRPDEAGRLPGEHLPQVPWDPGLTRS PSPRGPCRGAARAGHVGETPAPWGCPPPCAWEH KGPGSEGTP
3322	Α	1	420	AIVEDKHSGRSYDITSDLGNVLTSTSIAKTVNG*A ESSDSGAESDEEDAQEDLMGAYHSDIDKKMMKI VADHKNLEVIVTNGYDKDGFVHDIQNDIHASSSL NGRSTVHVKPIDENLGQTGKSAVCIHQDINDDH VEDVT
3323	A	8	459	DTLSLNCTLPETLPMTPSF*LSFL*FPGLARAKSIP TKTYSNEVVTLWYRPPDILLGSTDYSTQIDMW*G QVEVWQGPCGKGGGLVTTATQPAAFLFTVPSLP RGVGCIFYEMATGRPLFPGSTVEEQLHFIFRILSE EAWALCAVETHR
3324	A	1276	466	PGSTHASARITTY*L*IILSNATEVDNNFSKPPPFFP AGAPPASSSSSSSSSSSPTVSTAPPLIPPPGFPPPPG APPPSLIPTIESGHSSGYDSRSARAFPYGNVAFPH LPGSAPSWPSLVDTSKQWDYYARSSSSSSSSSSSS

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				SSSPRDRDRER*RTRERERERDHSPTPSVFNSDEE RYRYREYAERGYERHRASREKEERHRERRHREK EETRHKSSRSNSRRRHESEEGDSHRRHKHKKSKR SKEGKEAGSEPAPEQESTEATPAE
3325	A .	266	3312	TCLFSASCSSLPSPSSSFALLSTENTQRTYRVNPD GSLRVTFASGMEIGLSSEPHILAGAVNPTLGKCNI SLPGEHNANLISVL**GEQGCA*NVFHISFS*AHN RNLLSIDFDHITRTGKIYDDHRKFTLRILYDQTGR PILWSPVSRYNEVNITYSPSGLVTFIQRGTWNEK MEYDQSFL*SPQL*LSIICYSAFVSFQSVMLLLHS QRRYIFEYDQPDCLLSVTMPSMVRHSLQTMLSV GYYRNIYTPPDSSTSFIQDYSRDGRLLQTLHLGTG RRVLYKYTKQARLSEVLYDTTQVTLTYEESSGD LSDSSTLIA*LLTVFVLVPAGPLIGRQIFRFSEEGL VNARFDYSYNNFRVTSMQAVINETPLPIDLYRYV DVSGRTEQFGKFSVINYDLNQVITTTVMKHTKIF SANGQVIEVQYEILKAIAYWMTIQYDNVGRMVI CDIRVGVDANITRYFYEYDADGQLQTVSVNDKT QWRYSYDLNGNINLLSHGKSARLTPLRYDLRDRI TRLGEIQYKMDEDGFLRQRGNDIFEYNSNGLLQ KAYNKASGWTVQYYYDGLGRRVASKSSLGQHL QFFYADLTNPIRVTHLYNHTSSEITSLYYDLQGH LIAMELSSGEEYYVACDNTGTPLAVFSSRGQVIK EILYTPYGDIYHDTYPDFQVIIGFHGGLYDFLTKL VHLGQRDYDVVAGRWTTPNHHIWKQLNLLPKP FNLSTKLIKYGIFHFLFLILCLTDIRSWLELFGFQL HNVLPGFPKPELENSPSI*QMSNSMLHLLCASLS* TILGIQCELQKQLRNFISLDQLPMTPRYNDGRCLE GGKQPRFAAVPSVFGKGIKFAIKDGIVTADIIGVA NEDSRRLAAILNNAHYLENLHFTIEGRDTHYFIK LGSLEEDLVLIGNTGGRRILENGVNVTVSQMTSV LNGRTRRFADIQLQHGALCFNIRYGTTVEEEKNH VLEIARQRAVAQAWTKEQRRLQEGEEGIRAWTE GEKQQLLSTGRVQGYDGYFVLSVEQ
3326	A .	290	1041	KACLHLLSSFLTSNFLFNPLLPDSLYSVEARSQRA NLGPCRKKRLQTLMRLAAGFQYSSHKDPSLSAK EKHTDYHNEARGPWPGWVG*RTADGSCGRGPD GAHHPGPKSSSWRASRLLPGLGGSHHLDAYVGR DLECGTPAPLQLEIPPQPRGHPAPIPTGQAGPRDS GPGASP*VETRPLTDGRR*PGVRPVGWTPAHPAG TLRPRGAVEPSVSACGKWAPSPTSQGCCEGRCD AVPKHRAWRTPLCSQ
3327	A	1	418	CSECGKSFCKKSKFIIHQRTHTGEKPYECNQCGK SFCQKGTLTVHQRTHTGEKPYECNECGKNFYQK LHLIQHQRTHSGEKPYECSYCGKSFCQKTHLTQH QRTHSGERPYVCHDCGKTFSQKSALNDHQKIHT GVKLY
3328	A	1	270	VTRKLPIFIVDAFTARAFRGSPAADCLLENELDED MHQKIAREMNLSETAFIRKLHPTDNFAQRSCFGL IWFTPTTDLQILTSSILPSIL
3329	A	45	419	EELSCWQIWQQIANDLTRCQDSMINNSQCHKQG DFPYQVGTELSIQISEDENYIVNKADGPNNTGNP EFPILRTQDSWRKTFLTESQRLNRDQQISIKNKLC QCKKGVDPIGWISHHDGHRVHKR
3330	A	64	430	FWRNFTGLAPAAAVATTTSSSTMRFTSISNSLTST

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				AAIGLSFTTSTTTTATFTTNTTTTITSGFTVNQNQ LLSRGFENLVPYTSTVSVVTTPVMTYGHLEGLIN EGNLELEIKRRLSSQATQ
3331	A	3	407	TFGCSCTDCFFQKCCPAEAGVLLAYNKNQQIKIP PGTPIYECNSRCQCGPDCPNRIVQKGTQYSLCIFR TSNGRGWGVKTLVKIKRMSFVMEYVGEVITSEE AERRGQFYDNKGITYLFDLDYESDEFTVDAARY
3332	A	25	461	PAADFVLQARPTRADILGIHSKYDEVRKAGACFY KMTGLGPGPQALYNGEPFKHEEMNIKELKMAVL QRMMDASVYLQREVFLGTLNDRTNAIDFLMDR NNVVPRINTLILRTNQQYLNLLSTSVTADAEDFS TFFFLDSQDKSA
3333	A	317	54	AWIIFLPPLTSCPLWAPGTKHKTILEARSGLGPIK AYPRLGPPTPGEPEAPAQDRTFHCEICNVKVNSK VQLKQHISSRRHEIVDPV
3334	Α	304	410	AGPSLPSNLRQIFQSLPPFMDILLLLLFFMIIFAI
3335	A	19	418	VESRNSRVQPRVRLNDRTNAIDFLMDRNNVVPRI NTLILRTNQQYLNLISTSVTADVEDFSTFFFLDSQ DKSAVIAKNMYYLTQDDESIISAATLWIIADFDK PSGRKLLFNALKHMITSVHSRVGIIYNPFF
3336	A	1		PSSYSSDELSPGEPLTSPPWAPLGAPERPEHLLNR VLERLAGGATRDSAASDILLDDIVLTHSLFLPTEK FLQELHQYFVRAGGMEGPEGLGRKQACLAMLL HFLDTYQGLLQEEEGAGHIIKDLYLLIMKDESLY QGLREDTLRLHQLVETVELKIPEENQPPSKQVKP LFRHFRRIDSCLQTRVAFRGSDEIFCRVYMPDHS YVTIRSRLSASVQDILGSVTEKLQYSEEPAGREDS LILVAVSSSGEKVLLQPTEDCVFTALGINSHLFAC TRDSYEALVPLPEEIQVSPGDTEIHRVEPEDVANH LTAFHWELFRCVHELEFVDYVFHGE
3337	A	444	43	KILLCLANQFPDISFCPALPAVVALLLHYSIDEAE CFEKACRILACNDPGRRLIDQSFLAFESSCMTFGD LVNKYCQAAHKLMVAVSEDVLQVYADWQRWL FGELPLCYFARVFDVFLVEGYKVLYRVALAXXF
3338	A	1	398	FRGKVRGRSAEMPGSDTALTVDRTYSDPGRHHR CKSRVERHDMNTLSLPLNIRRGGSDTNLNFDVPD GILDFHKVKLTADSLKQKILKVTEQIKIEQTSRDG NVAEYLKLVNNADKQQAGRIKQVFEKKNQK
3339	A	1	665	AAAASNWGLITNIVNSIVGVSVLTMPFCFKQCGI VLGALLLVFCSWMTHQSCMFLVKSASLSKRRTY AGLAFHAYGKAGKMLVETSMIGLMLGTCIAFYV VIGDLGSNFFARLFGFQVGGTFRMFLLFAVSLCI VLPLSLQRNMMASIQSFSAMALLFYTVFMFVIVL SSLKHGLFSGQWLRRVSYVRWEGVFRCIPIFGMS FACQSQVLPTYDSLDEPSV
3340	A	198	367	LLPLQVLQEAFSRCVAVLTRSSKPSDMSVQVCG YISKCYSVAAQFEECREKITEMP
3341	A	562	277	HSVIKRTPRKYLAEIVLIDDFSNKEHLKEKLDEYI KLWNGLVKVFRNERREGLIQARSIGAQKAKLGQ VLIYLDAHCEVAVNWYAPLVAPISKDR
3342	A	385	2	NLTWWPLFRDVSFYIVDLIMLIIFFLDNVIMWWE SLLLLTAYFCYVVFMKFNVQVEKWVKQMINRN KVVKVTAPEAQAKPSAARDKDEPTLPAKPRLQR GGSSASLHNSLMRNSIFQNKIHTLDPHV
3343	Α	1	385	FRVDNSEEWKDVFIISSERSFKLDSLKCGTWYKV

SEQ ID NO:	Method	Predicted beginning nucleotide location corresponding to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion
				KLAAKNSVGSGRISEIIEAKTHGREPSFSKDQHLF THINSTHARLNLQGWNNGGCPITAIVLEYRPKGT WAWQGLRANSSGEVFLTELREATWY
3344	A	351	147	SPACITSSLSQHIADPRAAPTEVKVRVMNSTAISL QWNRVYSDTVQGQLREYRVRKPAPDSPNYPAH
3345	A	351	147	SPACITSSLSQHIADPRAAPTEVKVRVMNSTAISL QWNRVYSDTVQGQLREYRVRKPAPDSPNYPAH
3346	A	3	1509	AGIRHEAPPTTSNRHRRQIDRGVTHLNISGLKMP RGIAIDWVAGNVYWTDSGRDVIEVAQMKGENR KTLISGMIDEPHAIVVDPLRGTMYWSDWGNHPK IETAAMDGTLRETLVQDNIQWPTGLAVDYHNER LYWADAKLSVIGSIRLNGTDPIVAADSKRGLSHP FSIDVFEDYIYGVTYINNRVFKIHKFGHSPLVNLT GGLSHASDVVLYHQHKQPEVTNPCDRKKCEWL CLLSPSGPVCTCPNGKRLDNGTCVPVPSPTPPPD APRPGTCNLQCFNGGSCFLNARRQPKCRCQPRY TGDKCELDQCWEHCRNGGTCAASPSGMPTCRCP TGFTGPKCTQQVCAGYCANNSTCTVNQGNQPQ CRCLPGFLGDRCQYRQCSGYCENFGTCQMAAD GSRQCRCTAYFEGSRCEVNKCSRCLEGACVVNK QSGDVTCNCTDGRVAPSCLTCVGHCSNGGSCTM NSKMMPECQCPPHMTGPRCEEHVFSQQQPGHIA SILIP
3347	A	974	666	SPEMESHPITQAGVQWHHLSSLQPLPPGFK*FSCF SLPE*LGYRHVPPCLANSVFSVEMG\FLHVGQAG LELLTSGDLPALASQSAGITG\SHRARPENGFENIF
3348	A		1171	LSKITMPVICNEPLSFIQRLTEYM*HTYFIHRPSSL SDPVDRMQCVAAFAVSAVASQWERTGKPFNPLL GETYELVRDDLGFRLISEQVSHHPPISAFHAEGLN NDFIFHGSIYPKLKFWGKSVEAEPKGTITLELLEH NEAYTWTNPTCCVHNIIVGKLWIEQYGNVEIINH KTGDKCVLNFKPCGLFGKELHKVEGYIQDKSKK KLCALYGKWTECLYSVDPATFDAYKKNDKKNT EEKKNSKQMSTSEELDEMPVPDSESVFIIPGSVLL WRIAPRPPNSAQMYNFTSFAMVLNEVDKDMESV IPKTDCRLRPDIRAMENGEIDQASEEKKRLEEKQ RAARKNRSKSEEDWKTRWFHQGPNPYNGAQD WIYSGSYWDRNYFNLPDIY
3349	A	403	497	NFASSSGKYLRTQKIKCLNNKFTPFPTTEKK*SQS VRPP*SNRIY*ILQS*NISFS*LPN*NFASSSGKYLR TQKIKCLNNKFTPFPTTEKK
3350	A	1	712	GAPAQDCICLPFPFHSSFLESDIRKPARRKIQTTNP DFLLLLFMSVPVVSAPPFCPPAEGSRDGRPKASV ARPAAVHEHHSPRDCGHLPDVIRSSLGGWQPH*P AQPENRLL*LLPVE*GHQHPTVSPVP*AGSPGGAS GWPGPGQAWRVRVPGPHPLCPPASPPSPVQQ**E SVAAGSGLPGCVLCAAGRRPGPLPLLCVEVGQA LPPGAWVSSSGQRPGLTHPLAYSHGCVPSEG
3351	A	1	428	MAAVVAATALKGRGARNARVLRGILAGATANK ASHNRTRALQSHSSPEGKEEPEPLSPELEYIPRKR GKNPMKAVGLAWAIGFPCGILLFILTKREVDKDR VKQMKARQNMRLSNTGEYESQRFRASSQSAPSP DVGSGVQT
3352	A	2	841	RTLFRGRRRREDDRISRPHPSTAESKAPTPKFDLL ASNFPPLPGSSSRMPGELVLENRMSDVVKGVYK

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SEQ ID NO:	Method	Predicted beginning nucleotide location corresponding to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion
				EKDNEELTISCPVPADEQTECTSAQQLNMSTSSP CAAELTALSTTQQEKDLIEDSSVQKDGLNQTTIP VSPPSTTKPSRASTASPCNNNINAATAVALQEPR KLSYAEVCQKPPKEPSSVLVQPLRELRSNVVSPT KNEDNGAPENSVEKPHEKPEARASKDYSGFRGN IIPRGAAGKIREQRRQFSHRAIPQGVTRRNGKEQ YVPPRSPK
3353	A	1054	587	IATPTWTAPLTATPTPAHQYGPARVPNGAPRLEP PPGKRECRVGQYVVDLTSFEQLALPVLRNADCS SGPGQRVCVIDEIGKMELFSQLFIQAVRQTLSTPG TIILGTIPVPKGKPLALVEEIRNRKDVKVFNVTKE NRNHLLPDIVTCVQSSRK
3354	A	56	1268	GMEPVGCCGECRGSSVDPRSTFVLSNLAEVVER VLTFLPAKALLRVACVCRLWRECVRRVLRTHRS VTWISAGLAEAGHLEGHCLVRVVAEELENVRILP HTVLYMADSETFISLEECRGHKRARKRTSMETA LALEKLFPKQCQVLGIVTPGIVVTPMGSGSNRPQ EIEIGESGFALLFPQIEGIKIQPFHFIKDPKNLTLER HQLTEVGLLDNPELRVVLVFGYNCCKVGASNYL QVVSTFSDMNIILAGGQVDNLSSLTSEKNPLDI DASGVVGLSFSGHRIQSATVLLNEDVSDEKTAEA AMQRLKAANIPEHNTIGFMFACVGRGFQYYRAK GNVEADAFRKFFPSVPLFGFFGNGEIGCDRIVTG NFILRKCNEVKDDDLFHSYTTIMALIHLGSSK
3355	A	1	707	GTSSGLGGDRLAAPGPSPPSFYPQGRGERAYDIY SRLLRERIVCVMGPIDDSVASLVIAQLLFLQSESN KKPIHMYINSPGGVVTAGLAIYDTMQYILNPICT WCVGQAASMGSLLLAAGTPGMRHSLPNSRIMIH QPSGGARGQATDIAIQAEEIMKLKKQLYNIYAKH TKQSLQVIESAMERDRYMSPMEAQEFGILDKVL VHPPQDGEDEPTLVQKEPVEAAPAAEPVPAST
3356	A	352	338	FNYNFCRNLHMPSFLV*PGMCGLLAKHLSFHIVG AFLIT/LGVAALCKFAVA*PRKKAYADFYRNYN* IKEFEVRKANISQSTK
3357	A	1	403	ALGSCGGLLGTGLLKGTMSGTLWSKGIFAGYKR RIRIQREHTAVLKIEG\VYARDETEFYLRMICANV YKANNNTVTPVLTPDKTRVMWRKVTQAHGISI MVRAQFRTNLPADAIGHRIRMML*PSRMYTTEPS
3358	A	71	2897	FCSKDKCCLYLPDSINRSKSCTAKPGAHSQDRHA VMDSERQVKDTDDIESPKRSIRDSGYIDCWDSER SDSLSPPRHGRDDSFDSLDSFGSRSRQTPSPDVVL RGSSDGRGSDSESDLPHRKLPDVKKDDMSARRT SHGEPKSAVPFNQYLPNKSNQTAYVPAPLRKKK AEREEYRKSWSTATSPAGLGKKALQDYGPRTVPV S\DDAESTSMFDMRCEEEAAVQPHSRARQEQLQ LINNQLREEDDKWQDDLARWKSRKRSVSQDLIK KEEERKKMEKLLAGEDGTSERRKSIKTYREIVQE KERRERELHEAYKNARSQEEAEGILQQYIERFTIS EAVLERLEMPKILERSHSTEPNLSSFLNDPNPMK YLRQQSLPPPKFTATVETTIARASVLDTSMSAGS GSPSKTVTPKAVPMLTPKPYSQPKNSQDVLKTFK VDGKVSVNGETVHREEEKERECPTVAPAHSLTK SQMFEGVARVHGSPLELKQDNGSIEINIKKPNSV PQELAATTEKTEPNSQEDKNDGGKSRKGNIELAS SEPQHFTTTVTRCSPTVAFVEFPSSPQLKNDVSEE

SEQ ID NO:	Method	Predicted beginning nucleotide location corresponding to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion
				KDQKKPENEMSGKVELVLSQKVVKPKSPEPEAT LTFPFLDKMPEANQLHLPNLNSQVDSPSSEKSPV TTPFKFWAWDPEEERRRQEKWQQEQERLLQER YQ\KEQDK\LKEE\WEKAQKEVEEEERRYYEEEP* II\EDPVVPFTVSSSSADQLSTSSSMTEGSGTMNKI DLGNCQDEKQDRRWKKSFQGDDSDLLLKTRES DRLEEKGSLTEGALAHSGNPVSKGVHEDHQLDT EAGAPHCGTNPQLAQDPSQNQQTSNPTHSSEDV KPKTLPLDKSINHQIESPSERRKSISGKKLCSSCGL PLGKGAAMIIETLNLYFHIQCFRCG\ICKGQLGDA VSGTDVRIRNGLLNCNDCYMRSRSAGQPTTL
3359	A	3	368	EVTASREGRGACAWECGSSRGPWGLLRGTFAPV RAATP*S*LPKGSLRHRP*/CPPPVHLPPKSSCPPR AWAGRATSM*TSSYSSEYQPQTP*ALVTLPPRSY YLLTHLLTLTHLHHQILFEP
3360	A	2	392	ARGIGSLGRDHSGSGGGTGMAGAWVRKAADYV RSKDFRDYLMSTHFWGPVANWGLPIAAITDMK\ KSPEIISRRMTFAL*CYSLTFVRFAHYVQ\PWNWL MLGCHTAVDFDQLISSMPCISHGMTASASAL
3361		4619	532	LLLGRANSPPYNSVVRTLPPATLLLRRAGWESF WSCQSRSPWPPRPEVRAPAKGPRGVAGAAGACS AGARLGDAAGGDPASGQAARGCGARAPRGLGR TARARDTAMEDAGAAGPGPEPEPEPEPEPAPE PEPEPKPGAGTSEAFSRLWTDVMGILDGSLGNID DLAQQYADYYNTCFSDVCERMEELRKRRVSQD LEVEKPDASPTSLQLRSQIEESLGFCSAVSTPEVE RKNPLHKSNSEDSSVGKGDWKKKNKYFWQNFR KNQKGIMRQTSKGEDVGYVASEITMSDEERIQL MMMVKEKMITIEEALARLKEYEAQHRQSAALDP ADWPDGSYPTFDGSSNCNSREQSDDETEESVKF KRLHKLVNSTRRVRKKLIRVEEMKKPSTEGGEE HVFENSPVLDERSALYSGVHKKPLFFDGSPEKPP EDDSDSLTTSPSSSSLDTWGAGRKLVKTFSKGES RGLIKPPKKMGTFFSYPEEEKAQKVSRSLTEGEM KKGLGSLSHGRTCSFGGFDLTNRSLHVGSNNSDP MGKEGDFVYKEVIKSPTASRISLGKKVKSVKET MRKRMSKKYSSSVSEQDSGLDGMPGSPPPSQPD PEHLDKPKLKAGGSVESLRSSLSGQSSMSGQTVS TTDSSTSNRESVKSEDGDDEEPPYRGPFCGRARV HTDFTPSPYDTDSLKLKKGDIIDIISKPPMGTWMG LLNNKVGTFNFIYVDVLSED\EEKPKRPTRRRK GRPPQPKSVEDLLDRINLKEHMPTFLFNGYEDLD TFKLLEEEDLDELNIRDPEHRADLLTAVELLQEY DSNSDQSGSQEKLLVDSQGLSGCSPRDS*CYESS ENLENGKTRKASLLSAKSSTEPSLKAFSRNQLGN YPTLPLMKSGDALKQGQEEGRLGGGLAPDTSKS CDPPGC*LVLN\KNRRKPPSFPSCRSC\ETL\EGPQ TVDTWPRSHSLDDLQVEPGAEQDVPTEVTEPPPQ IVPEVPQKTTASSTKAQPLEQDSAVDNALLLTQS KRFSEPQKLTTKKLEGSIAASGRGLSPPQCLPRNY DAQPPGAKHGLARTPLEGHRKGHEFEGTHHPLG TKEGVDAEQRMQPKIPSQPPPVPAKKSRERLANG LHPVPMGPSGALPSPDAPCLPVKRGSPASPTSPSD CPPALAPRPLSGQALGSPPSTRPPPWLSELPENTS LQEHGVKLGPALTR\KVSCARGVDLETLTENKL\

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SEQ ID	Method	Predicted	Predicted end	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid,
NO:		beginning	nucleotide	E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine,
		nucleotide	location	I=Isoleucine, K=Lysine, L=Leucine, M=Methionine,
		location	corresponding to last amino	N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine,
		corresponding to first amino	acid residue of	X=Unknown, *=Stop codon, /=possible nucleotide deletion,
		acid residue of	peptide	\=possible nucleotide insertion
	}	peptide sequence	sequence	
		sequence		HAEGIRSSRREPYS*LRHGRCGI\P\EALVQRYAED
i				LDQPERDVAANMDQIRVKQLRKQHRMAIPSGGL
	}			TEICRKPVSPGCIS\SVSDWLISIGLPMYAGTLSTA
				GFSTL\SQVPSLSHTCLQEAG\ITEERHIRK\LLSAA
0060	ļ <u> </u>	 	4653	RLFKLPPGPEAM FRGGVGYAHTLHLLPFAGSSVVLARARRTDRWT
3362	A	1	4033	SGLVEMATLSLTVNSGDPPLGALLAVEHVKDDV
			Ì	SISVEEGKENILHVSENVIFTDVNSILRYLARVAT
Í				TAGLYGSNLMEHTEIDHWLEFSATKLSSCDSFTS
				TINELNHCLSLRTYLVGNSLSLADLCVWATLKG
		{		NAAWQEQLKQKKAPVHVKRWFGFLEAQQAFQS
		1		VGTKWDVSTTKARVAPEKKQDVGKFVELPGAE
			1	MGKVTVRFPPEASGYLHIGHAKAALLNQHYQV
				NFKGKLIMRFDDTNPEKEKEDFEKVILEDVAML
	ļ			HIKPDQFTYTSDHFETIMKYAEKLIQEGKAYVDD
				TPGEQIKAEREQRIESKHRKNPIEKNLQMWEEMK
				KGSQFGHSCCLRAKIDMSSNNGCMRDPTLYRCK
				IQPHPRTGN*Y\NV\YPTYDFACPIVDSIEGVTHAL
				RTTEYHDRDEQFYWIIEALGIRKPYIWEYSRLNL
		1	İ	NNTVLSKRKLTWFVNEGLVDGWDDPRFPTVRG
				VLRRGMTVEGLKQFIAAQGSSRSVVNMEWDKI
				WAFNKKVIDPVAPRYVALLKKEVIPVNVPEAQE
				EMKEVAKHPKNPEVGLKPVWYSPKVFIEGADAE
				TFSEGEMVTFINWGNLNITKIHKNADGKIISLDAK
				LNLENKDYKKTTKVTWLAETTHALPIPVICVTYE
				HLITKPVLGKDEDFKQYVNKNSKHEELMLGDPC
				LKDLKKGDIIQLQRRGFFICDQPYEPVSPYSCKEA
				PCVLIYIPDGHTKEMPTSGSKEKTKVEATKNETS
			}	APFKERPTPSLNNNCTTSEDSLVLYNRVAVQGD
			1	VVRELKAKKAPKEDVDAAVKQLLSLKAEYKEK
			1	TGQEYKPGNPPAEIGQNISSNSSASILESKSLYDE
				VAAQGEVVRKLKAEKSPKAKINEAVECLLSLKA
				QYKEKTGKEYIPGQPPLSQSSDSSPTRNSEPAGLE
			1	TPEAKVLFDKVASQGEVVRKLKTEKAPKDQVDI
				AVQELLQLKAQYKSLIGVEYKPVSATGAEDKDK
				KKKEKENKSEKQNKPQKQNDGQRKDPSKNQGG
	Į	1	}	GLSSSGAGEGQGPKKQTRLGLEAKK\EENLADW
	1			YSQVITKSEMIEYHDISGCYILRPWAYAIWEAIKD
				FFDAEIKKLGVENCYFPMFVSQSALEKEKTHVA
 	1			DFAPEVAWVTRSGKTELAEPIAIRPTSETVMYPA
	1	1	1	YAKWVQSHRDLPIKLNQWCNVVRWEFKHPQPF
į		1		LRTREFLWQEGHSAFATMEEAAEEVLQILDLYA
į	1	1		QVYEELLAIPVVKGRKTEKEKFAGGDYTTTIEAF
}		1		ISASGRAIQGGTSHHLGQNFSKMFEIVFEDPKIPG
]			EKQFAYQNSWGLTTRTIGVMTMVHGDNMGLVL
ł				PPRVACVQVVIIPCGITNALSEEDKEALIAKCNDY
		1	4	RRRLLSVNIRVRADLRDNYSPGWKFNHWELKG
		1		VPIRLEVGPRDMKSCQFVAVRRDTGEKLTVAEN
ı	1	1	1	EAETKLQAILEDIQVTLFTRASEDLKTHMVVANT
	1		1	MEDFQKILDSGKIVQIPFCGEIDCEDWIKKTTARD
			1	QDLEPGAPSMGAKSLCIPFKPLCELQPGAKCVCG
2262	 	2707	1614	KNPAKYYTLFGRSY
3363	Α	3797	1514	LGGAAPETMPFPVTTQGSQQTQPPQKHYGITSPIS
				LAAPKETDCVLTQK\LI\ETLKPFGGFLKKEEGTA
		<u></u>	<u> </u>	SRRNFNFGKN*INLVKEWIRRNQ*KAKNLPQSVI\

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SEQ ID NO:	Method	Predicted beginning nucleotide location corresponding	Predicted end nucleotide location corresponding to last amino	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine,
		to first amino acid residue of peptide sequence	acid residue of peptide sequence	X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion
				ENV\GGKIFT/FLGSYRL/GEVHTKGADIDGVCVF
	1			APRHVDRSDFFT\SFYDKLKLQEEVKDLRAVEEA FVPVIKLCFDGIEIDILFARLALQTIPEDLDLRDDS
				LLKNLDIRCIRSLNGCRVTDEILHLVPNIDNFRLT
			•	LRAIKLWAKRHNIYSNILGFLGGVSWAMLVART
				CQLYPNAIASTLVHKFFLVFSKWEWPNPVLLKQP
				EECNLNLPVWDPRVNPSDRYHLMPIITPAYPQQN
				STYNVSVSTRMVMVEEFKQGLAITDEILLSKAE WSKLFEAPNFFQKYKHYIVLLASAPTENQRLEW
				VGLVESKIRILVGSLEKNEFITLAHVNPQSFPAPK
				ENPDKEEFRTMWVIGLVFKKTENSENLSVDLTY
		ē		DIQSFTDTVYRQAINSKMFEVDMKIAAMHVKRK QLHQLLPNHVLQKKKKHSTEGVKLTALNDSSLD
				LSMDSDNSMSVPSPTSATKTSPLNSSGSSQGRNS
				PAPAVTAASVTNIQATEVSVPQVNSSESSGGTSSE
				SIPQTATQPAISPPPKPTVSRVVSSTRLVNPPPRSS
				GNAATSGNAATKIPTPIVGVKRTSSPHKEESPKK TKTEEDETSEDANCLALSGHDKTEAKEQLDTETS
				TTQSETIQTAASLLASQKTSSTDLSDIPALPANPIP
				VIKNSIKLRLNR
3364	Α	54	3073	SARTMSYDYHQNWGRDGGPRSSGGGYGGGPAG
	ļ			GHGGNRGSGGGGGGGGGGGGGWQGPASRAPER PRNRHVVREKTGAEEQ/WKRRGKREL/LVHMDE
				RREEQIVQLLNSVQAKNDKESEAQISWFAPEDHG
				YGTEVSTKNTPCSENKLDIQEKKLINQEKKMFRI
				RNRSYIDRDSEYLLQENEPDGTLDQKLLEDLQKK
				KNDLRYIEMQHFREKLPSYGMQKELVNLIDNHQ VTVISGETGCGKTTQVTQFILDNYIERGKGSACRI
]]	j	VCTQPRRISAISVAERVAAERAESCGSGNSTGYQI
				RLQSRLPRKQGSILYCTTGIILQWLQSDPYLSSVS
				HIVLDEIHERNLQSDVLMTVVKDLLNFRSDLKVI LMSATLNAEKFSEYFGNCPMIHIPGFTFPVVEYLL
				EDVIEKIRYVPEQKEHRCQFKRGFMQGHVNSQE
				KEEKEAIYKERWPDYVRELRRRYSASTVDVIEM
	:			MEDDKVDLNLIVALIRYIVLEEEDGAILVFLPGW
				DNISTLHDLLMSQVMFKSDKFLIIPLHSLMPTVN QTQVFKRTPPGVRKIVIATNIAETSITIDDVVYVID
				GGKIKETHFDTQNNISTMSAEWVSKANAKQRKG
				RAG\RVQPGSLLFICINGS*EASLLGWTIQLPEIF/R
				GTPLEELCLQIKVLRLGGI/GLFLSRLMDPPSNEA
				VLLSIRQL\RSLNALDKQEELTPLGVHLARLPVEP HIGKMILFGALFCCLDPVLTIAASLSFKDPFVIPLG
				KEKIADARRKELAKDTRSDHLTVVNAFEGWEEA
				RRRGFRYEKDYCWEYFLSSNTLQMLHNMKGQF
		•		AEHLLGAGFVSSRNPKDPESNINSDNEKIIKAVIC
				AGLYPKVAKIRLNLGKKRKMVKVYTKTDGLVA VHPKSVNVEQTDFHYNWLIYHLKMRTSSIYLYD
		1		CTEVSPYCLLFFGGDISIQKDNDQETIAVDEWIVF
				QSPARIAHLVKRAVVHMDERREEQIVQLLNSVQ
22.55		420	070	AKNDKESEAQISWFAPEDHGYDKKYFFKE
3365	Α	439	878	ECCNVRPLRETDLLKMKRKPRASSPVVEEQPRA NTKETRKKKSFSQPMSASTKEESQDGRRKGK*L
				KGRARKKNAPQKSMALRILEEGSRPTPSGHSDQL
				NEEL*QNELQLEQ/PEGT*LEQQSEGTQPEQQSGR
				MPTISTLSLSSE

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SEQ ID	Method	Predicted	Predicted end	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine,
NO:		beginning nucleotide	nucleotide location	I=Isoleucine, K=Lysine, L=Leucine, M=Methionine,
	1	location	corresponding	N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine,
	ļ	corresponding	to last amino	T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine,
		to first amino	acid residue of	X=Unknown, *=Stop codon, /=possible nucleotide deletion,
	1	acid residue of	peptide	\=possible nucleotide insertion
		peptide sequence	sequence	
3366	A	1	827	FRGYWGVREAFTDASWSGGLGPGKPGMKITRQ
3300	, A	1	027	KHAKKHLGFFRNNFGVREPYQILLDGTFCQAAL
		•		RGRIQLREQLPRYLMGETQLCTTRCVLKELETLG
				KDLYGAKLIAQKCQVRNCPHFKNAVSGSECLLS
				MVEEGNPHHYFVATQDQNLSVKVKKKPGVPLM
	ľ			FIIQNTMVLDKPSPKTIAFVKAVESG\RLSQCMRK
				KVSNISKRNRV**KTLNRGRRKKRKKISGPNPLS
				CLKKKKKAPDTQSSASEKKRKRKRIRNRSNPKV
				LSEKQNAEGE
3367	A	40	1467	MLWGCRAKACWGPRLSDLVASLSPQRECISVHV
İ			İ	GQAGVQIGNACWELFCLEHGIQADGTFDAQASK
				INDDDSFTTFFSETGNGKHVPRAVMIDLEPTVVD
		1		EVRAGTYRQLFHPEQLITGKEDAANNYARGHYT
į]			VGKESIDLVLDRIRKLTDACSGLQGFLIFHSFGGG
ł	ł	ľ	1	TGSGFTSLLMERLSLDYGKKSKLEFAIYPAPQVS
				TAVVEPYNSILTTHTTLEHSDCAFMVDNEAIYDI
				CRRNLDIERPTYTNLNRLISQIVSSITASLRFDGAL
				NVDLTEFQTNLVPYPRIHFPLVTYAPIISAEKAYH
			•	EQLSVAEITSSCFEPNSQMVKCDPRHGKYMACC
				MLYRGDVVPKDVNVAIAAIKTKRTIQFVDWCPT
				GFKVGINYQPPTVVPGGDLAKVQRAVCMLSNTT
		1		AIAEAWARLDHKFDLMYAKRAFVHWYVGEGM
				EEGEFS*RPGEDLA\ALE\KDYEEVGTDSFEEENE
				GEEF
3368	A	3	2597	SLLEETMDEDSSLREYTVSLDSDMDDASKCLQE
1 2200	1 1 2	1 3		
i	1			
				YDSGTGNTREALRPCPRTVSTKAQPGRSASSSSG
				YDSGTGNTREALRPCPRTVSTKAQPGRSASSSSG DKTTSFAEQKIRKLNHTDGESSGSSSQKTTPEGSE
				YDSGTGNTREALRPCPRTVSTKAQPGRSASSSSG DKTTSFAEQKIRKLNHTDGESSGSSSQKTTPEGSE LNIPHAGAWAQIPEETGLPQGRDTTQLLASEMV
				YDSGTGNTREALRPCPRTVSTKAQPGRSASSSSG DKTTSFAEQKIRKLNHTDGESSGSSSQKTTPEGSE LNIPHAGAWAQIPEETGLPQGRDTTQLLASEMV HLMMK\LKEKR\RAI*AQKKKMEAAFTKQRQKM
				YDSGTGNTREALRPCPRTVSTKAQPGRSASSSSG DKTTSFAEQKIRKLNHTDGESSGSSSQKTTPEGSE LNIPHAGAWAQIPEETGLPQGRDTTQLLASEMV HLMMK\LKEKR\RAI*AQKKKMEAAFTKQRQKM GRTAFLTVVKKKGDGISPLREEAAGAEDEKVYT
				YDSGTGNTREALRPCPRTVSTKAQPGRSASSSSG DKTTSFAEQKIRKLNHTDGESSGSSSQKTTPEGSE LNIPHAGAWAQIPEETGLPQGRDTTQLLASEMV HLMMK\LKEKR\RAI*AQKKKMEAAFTKQRQKM GRTAFLTVVKKKGDGISPLREEAAGAEDEKVYT DRAKEKESQKTDGQRSKSLADIKESMENPQAKW
				YDSGTGNTREALRPCPRTVSTKAQPGRSASSSSG DKTTSFAEQKIRKLNHTDGESSGSSSQKTTPEGSE LNIPHAGAWAQIPEETGLPQGRDTTQLLASEMV HLMMK\LKEKR\RAI*AQKKKMEAAFTKQRQKM GRTAFLTVVKKKGDGISPLREEAAGAEDEKVYT DRAKEKESQKTDGQRSKSLADIKESMENPQAKW LKSPTTPIDPEKQGNLASPSEETLNEGEILEYTKSI
·				YDSGTGNTREALRPCPRTVSTKAQPGRSASSSSG DKTTSFAEQKIRKLNHTDGESSGSSSQKTTPEGSE LNIPHAGAWAQIPEETGLPQGRDTTQLLASEMV HLMMK\LKEKR\RAI*AQKKKMEAAFTKQRQKM GRTAFLTVVKKKGDGISPLREEAAGAEDEKVYT DRAKEKESQKTDGQRSKSLADIKESMENPQAKW LKSPTTPIDPEKQGNLASPSEETLNEGEILEYTKSI EKLNSSLHFLQQEMQRLSLQQEMLMQMREQQS
·				YDSGTGNTREALRPCPRTVSTKAQPGRSASSSSG DKTTSFAEQKIRKLNHTDGESSGSSSQKTTPEGSE LNIPHAGAWAQIPEETGLPQGRDTTQLLASEMV HLMMK\LKEKR\RAI*AQKKKMEAAFTKQRQKM GRTAFLTVVKKKGDGISPLREEAAGAEDEKVYT DRAKEKESQKTDGQRSKSLADIKESMENPQAKW LKSPTTPIDPEKQGNLASPSEETLNEGEILEYTKSI EKLNSSLHFLQQEMQRLSLQQEMLMQMREQQS WVISPPQPSPQKQIRDFKPSKQAGLSSAIAPFSSD\
				YDSGTGNTREALRPCPRTVSTKAQPGRSASSSSG DKTTSFAEQKIRKLNHTDGESSGSSSQKTTPEGSE LNIPHAGAWAQIPEETGLPQGRDTTQLLASEMV HLMMK\LKEKR\RAI*AQKKKMEAAFTKQRQKM GRTAFLTVVKKKGDGISPLREEAAGAEDEKVYT DRAKEKESQKTDGQRSKSLADIKESMENPQAKW LKSPTTPIDPEKQGNLASPSEETLNEGEILEYTKSI EKLNSSLHFLQQEMQRLSLQQEMLMQMREQQS WVISPPQPSPQKQIRDFKPSKQAGLSSAIAPFSSD\ SPR\PTHPSSTSLLNRKSASFSVKSQRTPRPNELKI
				YDSGTGNTREALRPCPRTVSTKAQPGRSASSSG DKTTSFAEQKIRKLNHTDGESSGSSSQKTTPEGSE LNIPHAGAWAQIPEETGLPQGRDTTQLLASEMV HLMMK\LKEKR\RAI*AQKKKMEAAFTKQRQKM GRTAFLTVVKKKGDGISPLREEAAGAEDEKVYT DRAKEKESQKTDGQRSKSLADIKESMENPQAKW LKSPTTPIDPEKQGNLASPSEETLNEGEILEYTKSI EKLNSSLHFLQQEMQRLSLQQEMLMQMREQQS WVISPPQPSPQKQIRDFKPSKQAGLSSAIAPFSSD\ SPR\PTHPSSTSLLNRKSASFSVKSQRTPRPNELKI TPLNRTLTPPRSVDSLPRLRRFSPSQVPIQTRSFVC
				YDSGTGNTREALRPCPRTVSTKAQPGRSASSSSG DKTTSFAEQKIRKLNHTDGESSGSSSQKTTPEGSE LNIPHAGAWAQIPEETGLPQGRDTTQLLASEMV HLMMK\LKEKR\RAI*AQKKKMEAAFTKQRQKM GRTAFLTVVKKKGDGISPLREEAAGAEDEKVYT DRAKEKESQKTDGQRSKSLADIKESMENPQAKW LKSPTTPIDPEKQGNLASPSEETLNEGEILEYTKSI EKLNSSLHFLQQEMQRLSLQQEMLMQMREQQS WVISPPQPSPQKQIRDFKPSKQAGLSSAIAPFSSD\ SPR\PTHPSSTSLLNRKSASFSVKSQRTPRPNELKI TPLNRTLTPPRSVDSLPRLRRFSPSQVPIQTRSFVC FGDDGEPQLKESKPKEEVKKEELESKGTLEQRG
				YDSGTGNTREALRPCPRTVSTKAQPGRSASSSSG DKTTSFAEQKIRKLNHTDGESSGSSSQKTTPEGSE LNIPHAGAWAQIPEETGLPQGRDTTQLLASEMV HLMMK\LKEKR\RAI*AQKKKMEAAFTKQRQKM GRTAFLTVVKKKGDGISPLREEAAGAEDEKVYT DRAKEKESQKTDGQRSKSLADIKESMENPQAKW LKSPTTPIDPEKQGNLASPSEETLNEGEILEYTKSI EKLNSSLHFLQQEMQRLSLQQEMLMQMREQQS WVISPPQPSPQKQIRDFKPSKQAGLSSAIAPFSSD\ SPR\PTHPSSTSLLNRKSASFSVKSQRTPRPNELKI TPLNRTLTPPRSVDSLPRLRRFSPSQVPIQTRSFVC FGDDGEPQLKESKPKEEVKKEELESKGTLEQRG HNPEEKEIKPFESTVSEVLSLPVTETVCLTPNEDQ
				YDSGTGNTREALRPCPRTVSTKAQPGRSASSSSG DKTTSFAEQKIRKLNHTDGESSGSSSQKTTPEGSE LNIPHAGAWAQIPEETGLPQGRDTTQLLASEMV HLMMK\LKEKR\RAI*AQKKKMEAAFTKQRQKM GRTAFLTVVKKKGDGISPLREEAAGAEDEKVYT DRAKEKESQKTDGQRSKSLADIKESMENPQAKW LKSPTTPIDPEKQGNLASPSEETLNEGEILEYTKSI EKLNSSLHFLQQEMQRLSLQQEMLMQMREQQS WVISPPQPSPQKQIRDFKPSKQAGLSSAIAPFSSD\ SPR\PTHPSSTSLLNRKSASFSVKSQRTPRPNELKI TPLNRTLTPPRSVDSLPRLRRFSPSQVPIQTRSFVC FGDDGEPQLKESKPKEEVKKEELESKGTLEQRG HNPEEKEIKPFESTVSEVLSLPVTETVCLTPNEDQ LNQPTEPPPKPVFPPTAPKNVNLIEVSLSDLKPPE
				YDSGTGNTREALRPCPRTVSTKAQPGRSASSSG DKTTSFAEQKIRKLNHTDGESSGSSSQKTTPEGSE LNIPHAGAWAQIPEETGLPQGRDTTQLLASEMV HLMMK\LKEKR\RAI*AQKKKMEAAFTKQRQKM GRTAFLTVVKKKGDGISPLREEAAGAEDEKVYT DRAKEKESQKTDGQRSKSLADIKESMENPQAKW LKSPTTPIDPEKQGNLASPSEETLNEGEILEYTKSI EKLNSSLHFLQQEMQRLSLQQEMLMQMREQQS WVISPPQPSPQKQIRDFKPSKQAGLSSAIAPFSSD\ SPR\PTHPSSTSLLNRKSASFSVKSQRTPRPNELKI TPLNRTLTPPRSVDSLPRLRRFSPSQVPIQTRSFVC FGDDGEPQLKESKPKEEVKKEELESKGTLEQRG HNPEEKEIKPFESTVSEVLSLPVTETVCLTPNEDQ LNQPTEPPPKPVFPPTAPKNVNLIEVSLSDLKPPE KADVPVEKYDGESDKEQFDDDQKVCCGFFFKD
				YDSGTGNTREALRPCPRTVSTKAQPGRSASSSG DKTTSFAEQKIRKLNHTDGESSGSSSQKTTPEGSE LNIPHAGAWAQIPEETGLPQGRDTTQLLASEMV HLMMK\LKEKR\RAI*AQKKKMEAAFTKQRQKM GRTAFLTVVKKKGDGISPLREEAAGAEDEKVYT DRAKEKESQKTDGQRSKSLADIKESMENPQAKW LKSPTTPIDPEKQGNLASPSEETLNEGEILEYTKSI EKLNSSLHFLQQEMQRLSLQQEMLMQMREQQS WVISPPQPSPQKQIRDFKPSKQAGLSSAIAPFSSD\ SPR\PTHPSSTSLLNRKSASFSVKSQRTPRPNELKI TPLNRTLTPPRSVDSLPRLRRFSPSQVPIQTRSFVC FGDDGEPQLKESKPKEEVKKEELESKGTLEQRG HNPEEKEIKPFESTVSEVLSLPVTETVCLTPNEDQ LNQPTEPPPKPVFPPTAPKNVNLIEVSLSDLKPPE KADVPVEKYDGESDKEQFDDDQKVCCGFFFKD DQKAENDMAMKRAALLEKRLRREKETQLRKQQ
				YDSGTGNTREALRPCPRTVSTKAQPGRSASSSG DKTTSFAEQKIRKLNHTDGESSGSSSQKTTPEGSE LNIPHAGAWAQIPEETGLPQGRDTTQLLASEMV HLMMK\LKEKR\RAI*AQKKKMEAAFTKQRQKM GRTAFLTVVKKKGDGISPLREEAAGAEDEKVYT DRAKEKESQKTDGQRSKSLADIKESMENPQAKW LKSPTTPIDPEKQGNLASPSEETLNEGEILEYTKSI EKLNSSLHFLQQEMQRLSLQQEMLMQMREQQS WVISPPQPSPQKQIRDFKPSKQAGLSSAIAPFSSD\ SPR\PTHPSSTSLLNRKSASFSVKSQRTPRPNELKI TPLNRTLTPPRSVDSLPRLRRFSPSQVPIQTRSFVC FGDDGEPQLKESKPKEEVKKEELESKGTLEQRG HNPEEKEIKPFESTVSEVLSLPVTETVCLTPNEDQ LNQPTEPPPKPVFPPTAPKNVNLIEVSLSDLKPPE KADVPVEKYDGESDKEQFDDDQKVCCGFFFKD DQKAENDMAMKRAALLEKRLRREKETQLRKQQ LEAEMEHKKEETRRKTEEERQKKEDERARREFIR
				YDSGTGNTREALRPCPRTVSTKAQPGRSASSSG DKTTSFAEQKIRKLNHTDGESSGSSSQKTTPEGSE LNIPHAGAWAQIPEETGLPQGRDTTQLLASEMV HLMMK\LKEKR\RAI*AQKKKMEAAFTKQRQKM GRTAFLTVVKKKGDGISPLREEAAGAEDEKVYT DRAKEKESQKTDGQRSKSLADIKESMENPQAKW LKSPTTPIDPEKQGNLASPSEETLNEGEILEYTKSI EKLNSSLHFLQQEMQRLSLQQEMLMQMREQQS WVISPPQPSPQKQIRDFKPSKQAGLSSAIAPFSSD\ SPR\PTHPSSTSLLNRKSASFSVKSQRTPRPNELKI TPLNRTLTPPRSVDSLPRLRRFSPSQVPIQTRSFVC FGDDGEPQLKESKPKEEVKKEELESKGTLEQRG HNPEEKEIKPFESTVSEVLSLPVTETVCLTPNEDQ LNQPTEPPPKPVFPPTAPKNVNLIEVSLSDLKPPE KADVPVEKYDGESDKEQFDDDQKVCCGFFFKD DQKAENDMAMKRAALLEKRLRREKETQLRKQQ LEAEMEHKKEETRRKTEEERQKKEDERARREFIR QEYMRRKQLKLMEDMDTVIKPRPQVVKQKKQR
				YDSGTGNTREALRPCPRTVSTKAQPGRSASSSSG DKTTSFAEQKIRKLNHTDGESSGSSSQKTTPEGSE LNIPHAGAWAQIPEETGLPQGRDTTQLLASEMV HLMMK\LKEKR\RAI*AQKKKMEAAFTKQRQKM GRTAFLTVVKKKGDGISPLREEAAGAEDEKVYT DRAKEKESQKTDGQRSKSLADIKESMENPQAKW LKSPTTPIDPEKQGNLASPSEETLNEGEILEYTKSI EKLNSSLHFLQQEMQRLSLQQEMLMQMREQQS WVISPPQPSPQKQIRDFKPSKQAGLSSAIAPFSSD\ SPR\PTHPSSTSLLNRKSASFSVKSQRTPRPNELKI TPLNRTLTPPRSVDSLPRLRRFSPSQVPIQTRSFVC FGDDGEPQLKESKPKEEVKKEELESKGTLEQRG HNPEEKEIKPFESTVSEVLSLPVTETVCLTPNEDQ LNQPTEPPPKPVFPPTAPKNVNLIEVSLSDLKPPE KADVPVEKYDGESDKEQFDDDQKVCCGFFFKD DQKAENDMAMKRAALLEKRLRREKETQLRKQQ LEAEMEHKKEETRRKTEEERQKKEDERARREFIR QEYMRRKQLKLMEDMDTVIKPRPQVVKQKKQR PKSIHRDHIESPKTPIKGPPVSSLSLASLNTGDNES
				YDSGTGNTREALRPCPRTVSTKAQPGRSASSSSG DKTTSFAEQKIRKLNHTDGESSGSSSQKTTPEGSE LNIPHAGAWAQIPEETGLPQGRDTTQLLASEMV HLMMK\LKEKR\RAI*AQKKKMEAAFTKQRQKM GRTAFLTVVKKKGDGISPLREEAAGAEDEKVYT DRAKEKESQKTDGQRSKSLADIKESMENPQAKW LKSPTTPIDPEKQGNLASPSEETLNEGEILEYTKSI EKLNSSLHFLQQEMQRLSLQQEMLMQMREQQS WVISPPQPSPQKQIRDFKPSKQAGLSSAIAPFSSD\ SPR\PTHPSSTSLLNRKSASFSVKSQRTPRPNELKI TPLNRTLTPPRSVDSLPRLRRFSPSQVPIQTRSFVC FGDDGEPQLKESKPKEEVKKEELESKGTLEQRG HNPEEKEIKPFESTVSEVLSLPVTETVCLTPNEDQ LNQPTEPPPKPVFPPTAPKNVNLIEVSLSDLKPPE KADVPVEKYDGESDKEQFDDDQKVCCGFFFKD DQKAENDMAMKRAALLEKRLRREKETQLRKQQ LEAEMEHKKEETRRKTEEERQKKEDERARREFIR QEYMRRKQLKLMEDMDTVIKPRPQVVKQKKQR PKSIHRDHIESPKTPIKGPPVSSLSLASLNTGDNES VHSGKRTPRSESVEGFLSPSRCGSRNGEKDWEN
				YDSGTGNTREALRPCPRTVSTKAQPGRSASSSG DKTTSFAEQKIRKLNHTDGESSGSSSQKTTPEGSE LNIPHAGAWAQIPEETGLPQGRDTTQLLASEMV HLMMK\LKEKR\RAI*AQKKKMEAAFTKQRQKM GRTAFLTVVKKKGDGISPLREEAAGAEDEKVYT DRAKEKESQKTDGQRSKSLADIKESMENPQAKW LKSPTTPIDPEKQGNLASPSEETLNEGEILEYTKSI EKLNSSLHFLQQEMQRLSLQQEMLMQMREQQS WVISPPQPSPQKQIRDFKPSKQAGLSSAIAPFSSD\ SPR\PTHPSSTSLLNRKSASFSVKSQRTPRPNELKI TPLNRTLTPPRSVDSLPRLRRFSPSQVPIQTRSFVC FGDDGEPQLKESKPKEEVKKEELESKGTLEQRG HNPEEKEIKPFESTVSEVLSLPVTETVCLTPNEDQ LNQPTEPPPKPVFPPTAPKNVNLIEVSLSDLKPPE KADVPVEKYDGESDKEQFDDDQKVCCGFFFKD DQKAENDMAMKRAALLEKRLRREKETQLRKQQ LEAEMEHKKEETRRKTEEERQKKEDERARREFIR QEYMRRKQLKLMEDMDTVIKPRPQVVKQKKQR PKSIHRDHIESPKTPIKGPPVSSLSLASLNTGDNES VHSGKRTPRSESVEGFLSPSRCGSRNGEKDWEN ASTTSSVASGTEYTGPKLYKEPSAKSNKHIIQNAL
				YDSGTGNTREALRPCPRTVSTKAQPGRSASSSG DKTTSFAEQKIRKLNHTDGESSGSSSQKTTPEGSE LNIPHAGAWAQIPEETGLPQGRDTTQLLASEMV HLMMK\LKEKR\RAI*AQKKKMEAAFTKQRQKM GRTAFLTVVKKKGDGISPLREEAAGAEDEKVYT DRAKEKESQKTDGQRSKSLADIKESMENPQAKW LKSPTTPIDPEKQGNLASPSEETLNEGEILEYTKSI EKLNSSLHFLQQEMQRLSLQQEMLMQMREQQS WVISPPQPSPQKQIRDFKPSKQAGLSSAIAPFSSD\ SPR\PTHPSSTSLLNRKSASFSVKSQRTPRPNELKI TPLNRTLTPPRSVDSLPRLRRFSPSQVPIQTRSFVC FGDDGEPQLKESKPKEEVKKEELESKGTLEQRG HNPEEKEIKPFESTVSEVLSLPVTETVCLTPNEDQ LNQPTEPPPKPVFPPTAPKNVNLIEVSLSDLKPPE KADVPVEKYDGESDKEQFDDDQKVCCGFFFKD DQKAENDMAMKRAALLEKRLRREKETQLRKQQ LEAEMEHKKEETRRKTEEERQKKEDERARREFIR QEYMRRKQLKLMEDMDTVIKPRPQVVKQKKQR PKSIHRDHIESPKTPIKGPPVSSLSLASLNTGDNES VHSGKRTPRSESVEGFLSPSRCGSRNGEKDWEN ASTTSSVASGTEYTGPKLYKEPSAKSNKHIIQNAL AHCCLAGKVNEGQKKKILEEMEKSDANNFLILF
				YDSGTGNTREALRPCPRTVSTKAQPGRSASSSG DKTTSFAEQKIRKLNHTDGESSGSSSQKTTPEGSE LNIPHAGAWAQIPEETGLPQGRDTTQLLASEMV HLMMK\LKEKR\RAI*AQKKKMEAAFTKQRQKM GRTAFLTVVKKKGDGISPLREEAAGAEDEKVYT DRAKEKESQKTDGQRSKSLADIKESMENPQAKW LKSPTTPIDPEKQGNLASPSEETLNEGEILEYTKSI EKLNSSLHFLQQEMQRLSLQQEMLMQMREQQS WVISPPQPSPQKQIRDFKPSKQAGLSSAIAPFSSD\ SPR\PTHPSSTSLLNRKSASFSVKSQRTPRPNELKI TPLNRTLTPPRSVDSLPRLRRFSPSQVPIQTRSFVC FGDDGEPQLKESKPKEEVKKEELESKGTLEQRG HNPEEKEIKPFESTVSEVLSLPVTETVCLTPNEDQ LNQPTEPPPKPVFPPTAPKNVNLIEVSLSDLKPPE KADVPVEKYDGESDKEQFDDDQKVCCGFFFKD DQKAENDMAMKRAALLEKRLRREKETQLRKQQ LEAEMEHKKEETRRKTEEERQKKEDERARREFIR QEYMRRKQLKLMEDMDTVIKPRPQVVKQKKQR PKSIHRDHIESPKTPIKGPPVSSLSLASLNTGDNES VHSGKRTPRSESVEGFLSPSRCGSRNGEKDWEN ASTTSSVASGTEYTGPKLYKEPSAKSNKHIIQNAL AHCCLAGKVNEGQKKKILEEMEKSDANNFLILF RDSGCQFRSLYTYCPETEEINKLTGIGPKSITKKM
				YDSGTGNTREALRPCPRTVSTKAQPGRSASSSG DKTTSFAEQKIRKLNHTDGESSGSSSQKTTPEGSE LNIPHAGAWAQIPEETGLPQGRDTTQLLASEMV HLMMK\LKEKR\RAI*AQKKKMEAAFTKQRQKM GRTAFLTVVKKKGDGISPLREEAAGAEDEKVYT DRAKEKESQKTDGQRSKSLADIKESMENPQAKW LKSPTTPIDPEKQGNLASPSEETLNEGEILEYTKSI EKLNSSLHFLQQEMQRLSLQQEMLMQMREQQS WVISPPQPSPQKQIRDFKPSKQAGLSSAIAPFSSD\ SPR\PTHPSSTSLLNRKSASFSVKSQRTPRPNELKI TPLNRTLTPPRSVDSLPRLRRFSPSQVPIQTRSFVC FGDDGEPQLKESKPKEEVKKEELESKGTLEQRG HNPEEKEIKPFESTVSEVLSLPVTETVCLTPNEDQ LNQPTEPPPKPVFPPTAPKNVNLIEVSLSDLKPPE KADVPVEKYDGESDKEQFDDDQKVCCGFFFKD DQKAENDMAMKRAALLEKRLRREKETQLRKQQ LEAEMEHKKEETRRKTEEERQKKEDERARREFIR QEYMRRKQLKLMEDMDTVIKPRPQVVKQKKQR PKSIHRDHIESPKTPIKGPPVSSLSLASLNTGDNES VHSGKRTPRSESVEGFLSPSRCGSRNGEKDWEN ASTTSSVASGTEYTGPKLYKEPSAKSNKHIIQNAL AHCCLAGKVNEGQKKKILEEMEKSDANNFLILF
				YDSGTGNTREALRPCPRTVSTKAQPGRSASSSG DKTTSFAEQKIRKLNHTDGESSGSSSQKTTPEGSE LNIPHAGAWAQIPEETGLPQGRDTTQLLASEMV HLMMK\LKEKR\RAI*AQKKKMEAAFTKQRQKM GRTAFLTVVKKKGDGISPLREEAAGAEDEKVYT DRAKEKESQKTDGQRSKSLADIKESMENPQAKW LKSPTTPIDPEKQGNLASPSEETLNEGEILEYTKSI EKLNSSLHFLQQEMQRLSLQQEMLMQMREQQS WVISPPQPSPQKQIRDFKPSKQAGLSSAIAPFSSD\ SPR\PTHPSSTSLLNRKSASFSVKSQRTPRPNELKI TPLNRTLTPPRSVDSLPRLRRFSPSQVPIQTRSFVC FGDDGEPQLKESKPKEEVKKEELESKGTLEQRG HNPEEKEIKPFESTVSEVLSLPVTETVCLTPNEDQ LNQPTEPPPKPVFPPTAPKNVNLIEVSLSDLKPPE KADVPVEKYDGESDKEQFDDDQKVCCGFFFKD DQKAENDMAMKRAALLEKRLRREKETQLRKQQ LEAEMEHKKEETRRKTEEERQKKEDERARREFIR QEYMRRKQLKLMEDMDTVIKPRPQVVKQKKQR PKSIHRDHIESPKTPIKGPPVSSLSLASLNTGDNES VHSGKRTPRSESVEGFLSPSRCGSRNGEKDWEN ASTTSSVASGTEYTGPKLYKEPSAKSNKHIIQNAL AHCCLAGKVNEGQKKKILEEMEKSDANNFLILF RDSGCQFRSLYTYCPETEEINKLTGIGPKSITKKM
3360	A	977	594	YDSGTGNTREALRPCPRTVSTKAQPGRSASSSSG DKTTSFAEQKIRKLNHTDGESSGSSSQKTTPEGSE LNIPHAGAWAQIPEETGLPQGRDTTQLLASEMV HLMMK\LKEKR\RAI*AQKKKMEAAFTKQRQKM GRTAFLTVVKKKGDGISPLREEAAGAEDEKVYT DRAKEKESQKTDGQRSKSLADIKESMENPQAKW LKSPTTPIDPEKQGNLASPSEETLNEGEILEYTKSI EKLNSSLHFLQQEMQRLSLQQEMLMQMREQQS WVISPPQPSPQKQIRDFKPSKQAGLSSAIAPFSSD\ SPR\PTHPSSTSLLNRKSASFSVKSQRTPRPNELKI TPLNRTLTPPRSVDSLPRLRRFSPSQVPIQTRSFVC FGDDGEPQLKESKPKEEVKKEELESKGTLEQRG HNPEEKEIKPFESTVSEVLSLPVTETVCLTPNEDQ LNQPTEPPPKPVFPPTAPKNVNLIEVSLSDLKPPE KADVPVEKYDGESDKEQFDDDQKVCCGFFFKD DQKAENDMAMKRAALLEKRLRREKETQLRKQQ LEAEMEHKKEETRRKTEEERQKKEDERARREFIR QEYMRRKQLKLMEDMDTVIKPRPQVVKQKKQR PKSIHRDHIESPKTPIKGPPVSSLSLASLNTGDNES VHSGKRTPRSESVEGFLSPSRCGSRNGEKDWEN ASTTSSVASGTEYTGPKLYKEPSAKSNKHIIQNAL AHCCLAGKVNEGQKKKILEEMEKSDANNFLILF RDSGCQFRSLYTYCPETEEINKLTGIGPKSITKKM IEGLYKYNSDRKQFSHIPAKTLSASVDAITIHSHL WQTKRPVTPKKLLPTKA
3369	A	977	594	YDSGTGNTREALRPCPRTVSTKAQPGRSASSSSG DKTTSFAEQKIRKLNHTDGESSGSSSQKTTPEGSE LNIPHAGAWAQIPEETGLPQGRDTTQLLASEMV HLMMK\LKEKR\RAI*AQKKKMEAAFTKQRQKM GRTAFLTVVKKKGDGISPLREEAAGAEDEKVYT DRAKEKESQKTDGQRSKSLADIKESMENPQAKW LKSPTTPIDPEKQGNLASPSEETLNEGEILEYTKSI EKLNSSLHFLQQEMQRLSLQQEMLMQMREQQS WVISPPQPSPQKQIRDFKPSKQAGLSSAIAPFSSD\ SPR\PTHPSSTSLLNRKSASFSVKSQRTPRPNELKI TPLNRTLTPPRSVDSLPRLRRFSPSQVPIQTRSFVC FGDDGEPQLKESKPKEEVKKEELESKGTLEQRG HNPEEKEIKPFESTVSEVLSLPVTETVCLTPNEDQ LNQPTEPPPKPVFPPTAPKNVNLIEVSLSDLKPPE KADVPVEKYDGESDKEQFDDDQKVCCGFFFKD DQKAENDMAMKRAALLEKRLRREKETQLRKQQ LEAEMEHKKEETRRKTEEERQKKEDERARREFIR QEYMRRKQLKLMEDMDTVIKPRPQVVKQKKQR PKSIHRDHIESPKTPIKGPPVSSLSLASLNTGDNES VHSGKRTPRSESVEGFLSPSRCGSRNGEKDWEN ASTTSSVASGTEYTGPKLYKEPSAKSNKHIIQNAL AHCCLAGKVNEGQKKKILEEMEKSDANNFLILF RDSGCQFRSLYTYCPETEEINKLTGIGPKSITKKM IEGLYKYNSDRKQFSHIPAKTLSASVDAITIHSHL WQTKRPVTPKKLLPTKA
3369	A	977	594	YDSGTGNTREALRPCPRTVSTKAQPGRSASSSSG DKTTSFAEQKIRKLNHTDGESSGSSSQKTTPEGSE LNIPHAGAWAQIPEETGLPQGRDTTQLLASEMV HLMMK\LKEKR\RAI*AQKKKMEAAFTKQRQKM GRTAFLTVVKKKGDGISPLREEAAGAEDEKVYT DRAKEKESQKTDGQRSKSLADIKESMENPQAKW LKSPTTPIDPEKQGNLASPSEETLNEGEILEYTKSI EKLNSSLHFLQQEMQRLSLQQEMLMQMREQQS WVISPPQPSPQKQIRDFKPSKQAGLSSAIAPFSSD\ SPR\PTHPSSTSLLNRKSASFSVKSQRTPRPNELKI TPLNRTLTPPRSVDSLPRLRRFSPSQVPIQTRSFVC FGDDGEPQLKESKPKEEVKKEELESKGTLEQRG HNPEEKEIKPFESTVSEVLSLPVTETVCLTPNEDQ LNQPTEPPPKPVFPPTAPKNVNLIEVSLSDLKPPE KADVPVEKYDGESDKEQFDDDQKVCCGFFFKD DQKAENDMAMKRAALLEKRLRREKETQLRKQQ LEAEMEHKKEETRRKTEEERQKKEDERARREFIR QEYMRRKQLKLMEDMDTVIKPRPQVVKQKKQR PKSIHRDHIESPKTPIKGPPVSSLSLASLNTGDNES VHSGKRTPRSESVEGFLSPSRCGSRNGEKDWEN ASTTSSVASGTEYTGPKLYKEPSAKSNKHIIQNAL AHCCLAGKVNEGQKKKILEEMEKSDANNFLILF RDSGCQFRSLYTYCPETEEINKLTGIGPKSITKKM IEGLYKYNSDRKQFSHIPAKTLSASVDAITIHSHL WQTKRPVTPKKLLPTKA RGSGLTQEPGSVGQLALACAEGAVEWLYPAGAL RLTLGGPDPRARPGIACLRPVRPFAGAQVFAERA
3369	A	977	594	YDSGTGNTREALRPCPRTVSTKAQPGRSASSSG DKTTSFAEQKIRKLNHTDGESSGSSSQKTTPEGSE LNIPHAGAWAQIPEETGLPQGRDTTQLLASEMV HLMMK\LKEKR\RAI*AQKKKMEAAFTKQRQKM GRTAFLTVVKKKGDGISPLREEAAGAEDEKVYT DRAKEKESQKTDGQRSKSLADIKESMENPQAKW LKSPTTPIDPEKQGNLASPSEETLNEGEILEYTKSI EKLNSSLHFLQQEMQRLSLQQEMLMQMREQQS WVISPPQPSPQKQIRDFKPSKQAGLSSAIAPFSSD\ SPR\PTHPSSTSLLNRKSASFSVKSQRTPRPNELKI TPLNRTLTPPRSVDSLPRLRRFSPSQVPIQTRSFVC FGDDGEPQLKESKPKEEVKKEELESKGTLEQRG HNPEEKEIKPFESTVSEVLSLPVTETVCLTPNEDQ LNQPTEPPPKPVFPPTAPKNVNLIEVSLSDLKPPE KADVPVEKYDGESDKEQFDDDQKVCCGFFFKD DQKAENDMAMKRAALLEKRLRREKETQLRKQQ LEAEMEHKKEETRRKTEEERQKKEDERARREFIR QEYMRRKQLKLMEDMDTVIKPRPQVVKQKKQR PKSIHRDHIESPKTPIKGPPVSSLSLASLNTGDNES VHSGKRTPRSESVEGFLSPSRCGSRNGEKDWEN ASTTSSVASGTEYTGPKLYKEPSAKSNKHIIQNAL AHCCLAGKVNEGQKKKILEEMEKSDANNFLILF RDSGCQFRSLYTYCPETEEINKLTGIGPKSITKKM IEGLYKYNSDRKQFSHIPAKTLSASVDAITIHSHL WQTKRPVTPKKLLPTKA RGSGLTQEPGSVGQLALACAEGAVEWLYPAGAL RLTLGGPDPRARPGIACLRPVRPFAGAQVFAERA GGALELLLAEGPGPAGGRCVRWGPRERRALFLQ
3369	A	977	594	YDSGTGNTREALRPCPRTVSTKAQPGRSASSSSG DKTTSFAEQKIRKLNHTDGESSGSSSQKTTPEGSE LNIPHAGAWAQIPEETGLPQGRDTTQLLASEMV HLMMK\LKEKR\RAI*AQKKKMEAAFTKQRQKM GRTAFLTVVKKKGDGISPLREEAAGAEDEKVYT DRAKEKESQKTDGQRSKSLADIKESMENPQAKW LKSPTTPIDPEKQGNLASPSEETLNEGEILEYTKSI EKLNSSLHFLQQEMQRLSLQQEMLMQMREQQS WVISPPQPSPQKQIRDFKPSKQAGLSSAIAPFSSD\ SPR\PTHPSSTSLLNRKSASFSVKSQRTPRPNELKI TPLNRTLTPPRSVDSLPRLRRFSPSQVPIQTRSFVC FGDDGEPQLKESKPKEEVKKEELESKGTLEQRG HNPEEKEIKPFESTVSEVLSLPVTETVCLTPNEDQ LNQPTEPPPKPVFPPTAPKNVNLIEVSLSDLKPPE KADVPVEKYDGESDKEQFDDDQKVCCGFFFKD DQKAENDMAMKRAALLEKRLRREKETQLRKQQ LEAEMEHKKEETRRKTEEERQKKEDERARREFIR QEYMRRKQLKLMEDMDTVIKPRPQVVKQKKQR PKSIHRDHIESPKTPIKGPPVSSLSLASLNTGDNES VHSGKRTPRSESVEGFLSPSRCGSRNGEKDWEN ASTTSSVASGTEYTGPKLYKEPSAKSNKHIIQNAL AHCCLAGKVNEGQKKKILEEMEKSDANNFLILF RDSGCQFRSLYTYCPETEEINKLTGIGPKSITKKM IEGLYKYNSDRKQFSHIPAKTLSASVDAITIHSHL WQTKRPVTPKKLLPTKA RGSGLTQEPGSVGQLALACAEGAVEWLYPAGAL RLTLGGPDPRARPGIACLRPVRPFAGAQVFAERA

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A U A1 U D U A1 U 1 U 2 U

SEQ ID NO: Method Predicted beginning mucleotide location corresponding corresponding to first amino acid residue of peptide sequence sequence Seque	listidine, e, S≒Serine, e,
Invalential location corresponding to first amino acid residue of peptide sequence International content o	e, S=Serine, e,
location corresponding to first amino acid residue of peptide sequence Time content of peptide sequence Time conten	e,
to first amino acid residue of peptide sequence X=Unknown, *=Stop codon, =possible nucleotide de peptide sequence	e, etion
acid residue of peptide sequence Sequence Sequence Sequence Sequence Sequence	etion
Peptide sequence YSAVLFPC*AMDHLESFIAECDRRTELA TQEEISAEVSAKAEKVHELNEEIGKLLAI AEGNVDESQKILMEVEKVRAKKKEAEK QEKRNQDRLRRREERERELSRRSGSB SRSRDRRRRSRSTSRERKLSRSSRDI) SRSRSHSRGHRRASRDRSAKYKFSREA ESGRSERGPPDWRLESSNGKMASRRSEI DLLNRMIVWKHGLLI 3371 A 345 1383 DLSLECTGFKETNLGVYFLSSKWVLRLY YSAVLFPC*AMDHLESFIAECDRRTELA TQEEISAEVSAKAEKVHELNEEIGKLLAI AEGNVDESQKILMEVEKVRAKKKEAEK QEKRNQDRLRRREEREREERLSRRSGSB SRSRDRRRRSSTSRERKLSRSRSRDI SRSRSHSRGHRRASRDRSAKYKFSREAE ESGRSERGPPDWRLESSNGKMASRRSEI DLLNRMIVWKHGLLI 3372 A 239 3348 PMQNCMCSLTLSVLPLGPQPPVPEKRPP MSDDVHSLGKVTSDLAKRRKLTS**SGI ARRSGEVTLTKGDPGSLEEWETVVGDD SYSVDERVDSDSKSEVEALTEQLSEEEE EEEEEEEEEEEEEEEEDEESGNQSDRSGSSGI KWRKDSPWVKPSRKRRKEPPRAKEPR GSSGPSEYMEVPLGSLELPSEGTLSPNH TSSLETERGFEELPLCSCRMEAPKIDRISI CMATESVDGELSGCNAAILKRETMRPS: VLCETHRARMVKHHCCPGCGYFCTAG*	cuou,
YSAVLFPC*AMDHLESFIAECDRRTELA TQEEISAEVSAKAEKVHELNEEIGKLLAI AEGNVDESQKILMEVEKVRAKKKEAEK QEKRNQDRLRRREERERERLSRRSGSR SRSRDRRRRSRSTSRERRKLSRSRSRDI SRSRSHSRGHRRASRDRSAKYKFSRERA ESGRSERGPPDWRLESSNGKMASRRSEI DLLNRMIVWKHGLLI 3371 A 345 1383 DLSLECTGFKETINLGVYFLSSKWVLRLY YSAVLFPC*AMDHLESFIAECDRRTELA TQEEISAEVSAKAEKVHELNEEIGKLLAI AEGNVDESQKILMEVEKVRAKKKEAEK QEKRNQDRLRRREEREREERLSRRSGSF SRSRDRRRRSRSTSRERKLSRSRSDI SRSRSHSRGHRRASRDRSAKYKFSRERA ESGRSERGPPDWRLESSNGKMASRRSEI DLLNRMIVWKHGLLI 3372 A 239 3348 PMQNCMCSLTLSVLPLGPQPPVPEKRPP MSDDVHSLGKVTSDLAKRRKLTS*GGI ARRSGEVTLTKGDPGSLEEWETVVGDD SYSVDERVDSDSKSEVEALTEQLSEEEE EEEEEEEEEEEEEEDEESGNQSDRSGSSGI KWRKDSPWVKPSRKRRKREPPRAKEPR GSSGPSEYMEVPLGSLELPSEGTLSPNH TSSLETERGFEELPLCSCRMEAPKIDRISI CMATESVDGELSGCNAAILKRETMRPSI VLCETHRARMVKHHCCPGCGYFCTAG*	
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QEKRNQDRLRRREERERLSRRSGSR SRSRDRRRRSRSTSRERRKLSRSRSRDI SRSRSHSRGHRRASRDRSAKYKFSRERA ESGRSERGPPDWRLESSNGKMASRRSEI DLLNRMIVWKHGLLI 3371 A 345 1383 DLSLECTGFKETNLGVYFLSSKWVLRLY YSAVLFPC*AMDHLESFIAECDRRTELA TQEEISAEVSAKAEKVHELNEEIGKLLA: AEGNVDESQKILMEVEKVRAKKKEAEK QEKRNQDRLRRREEREREERLSRRSGSR SRSRDRRRRSRSTSRERRKLSRSRSDI SRSRSHSRGHRRASRDRSAKYKFSRERA ESGRSERGPPDWRLESSNGKMASRRSEI DLLNRMIVWKHGLLI 3372 A 239 3348 PMQNCMCSLTLSVLPLGPQPPVPEKRPP MSDDVHSLGKVTSDLAKRRKLTS*GGI ARRSGEVTLTKGDPGSLEEWETVVGDD SYSVDERVDSDSKSEVEALTEQLSEEEE EEEEEEEEEEEEEEEEEEEEEEGRQSDRSGSSGI KWRDSPWVKPSRKRRKREPPRAKEPR GSSGPSEYMEVPLGSLELPSEGTLSPNHL TSSLETERGFEELPLCSCRMEAPKIDRISI CMATESVDGELSGCNAAILKRETMRPS VLCETHRARMVKHHCCPGCGYFCTAG	KAEQLG
QEKRNQDRLRRREERERLSRRSGSR SRSRDRRRRSRSTSRERRKLSRSRSRDI SRSRSHSRGHRRASRDRSAKYKFSRERA ESGRSERGPPDWRLESSNGKMASRRSEI DLLNRMIVWKHGLLI 3371 A 345 1383 DLSLECTGFKETNLGVYFLSSKWVLRLY YSAVLFPC*AMDHLESFIAECDRRTELA TQEEISAEVSAKAEKVHELNEEIGKLLA: AEGNVDESQKILMEVEKVRAKKKEAEK QEKRNQDRLRRREEREREERLSRRSGSR SRSRDRRRRSRSTSRERRKLSRSRSDI SRSRSHSRGHRRASRDRSAKYKFSRERA ESGRSERGPPDWRLESSNGKMASRRSEI DLLNRMIVWKHGLLI 3372 A 239 3348 PMQNCMCSLTLSVLPLGPQPPVPEKRPP MSDDVHSLGKVTSDLAKRRKLTS*GGI ARRSGEVTLTKGDPGSLEEWETVVGDD SYSVDERVDSDSKSEVEALTEQLSEEEE EEEEEEEEEEEEEEEEEEEEEEGRQSDRSGSSGI KWRDSPWVKPSRKRRKREPPRAKEPR GSSGPSEYMEVPLGSLELPSEGTLSPNHL TSSLETERGFEELPLCSCRMEAPKIDRISI CMATESVDGELSGCNAAILKRETMRPS VLCETHRARMVKHHCCPGCGYFCTAG	TVAEK
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TQEEISAEVSAKAEKVHELNEEIGKLLA AEGNVDESQKILMEVEKVRAKKKEAEK QEKRNQDRLRRREERERERLSRRSGSE SRSRDRRRRRSSTSRERRKLSRSRSRDI SRSRSHSGHRRASRDRSAKYKFSRERA ESGRSERGPPDWRLESSNGKMASRRSEI DLLNRMIVWKHGLLI 3372 A 239 3348 PMQNCMCSLTLSVLPLGPQPPVPEKRPP MSDDVHSLGKVTSDLAKRRKLTS*GGI ARRSGEVTLTKGDPGSLEEWETVVGDD SYSVDERVDSDSKSEVEALTEQLSEEEE EEEEEEEEEEEEEEEEDEESGNQSDRSGSSGI KWRKDSPWVKPSRKRRKREPPRAKEPR GSSGPSEYMEVPLGSLELPSEGTLSPNHL TSSLETERGFEELPLCSCRMEAPKIDRISI CMATESVDGELSGCNAAILKRETMRPS VLCETHRARMVKHHCCPGCGYFCTAG	
AEGNVDESQKILMEVEKVRAKKKEAEK QEKRNQDRLRRREERERERLSRRSGSF SRSRDRRRRRSRSTSRERRKLSRSRSRDI SRSRSHSRGHRRASRDRSAKYKFSRERA ESGRSERGPPDWRLESSNGKMASRRSEI DLLNRMIVWKHGLLI 3372 A 239 3348 PMQNCMCSLTLSVLPLGPQPPVPEKRPP MSDDVHSLGKVTSDLAKRRKLTS*GGI ARRSGEVTLTKGDPGSLEEWETVVGDD SYSVDERVDSDSKSEVEALTEQLSEEEE EEEEEEEEEEEEEEEEEEESGNQSDRSGSSGI KWRKDSPWVKPSRKRRKREPPRAKEPR GSSGPSEYMEVPLGSLELPSEGTLSPNHL TSSLETERGFEELPLCSCRMEAPKIDRISI CMATESVDGELSGCNAAILKRETMRPS VLCETHRARMVKHHCCPGCGYFCTAG	
QEKRNQDRLRRREEREERLSRRSGSR SRSRDRRRRRSRSTSRERRKLSRSRSRDI SRSRSHSRGHRRASRDRSAKYKFSRERA ESGRSERGPPDWRLESSNGKMASRRSEI DLLNRMIVWKHGLLI 3372 A 239 3348 PMQNCMCSLTLSVLPLGPQPPVPEKRPP MSDDVHSLGKVTSDLAKRRKLTS*GGI ARRSGEVTLTKGDPGSLEEWETVVGDD SYSVDERVDSDSKSEVEALTEQLSEEEE EEEEEEEEEEEEEEEEEDEESGNQSDRSGSSGI KWRKDSPWVKPSRKRRKREPPRAKEPR GSSGPSEYMEVPLGSLELPSEGTLSPNHL TSSLETERGFEELPLCSCRMEAPKIDRISI CMATESVDGELSGCNAAILKRETMRPSS VLCETHRARMVKHHCCPGCGYFCTAG	
SRSRDRRRRSSTSRERRKLSRSRSRDD SRSRSHSRGHRRASRDRSAKYKFSRERA ESGRSERGPPDWRLESSNGKMASRRSEI DLLNRMIVWKHGLLI 3372 A 239 3348 PMQNCMCSLTLSVLPLGPQPPVPEKRPP MSDDVHSLGKVTSDLAKRRKLTS*GGI ARRSGEVTLTKGDPGSLEEWETVVGDD SYSVDERVDSDSKSEVEALTEQLSEEEE EEEEEEEEEEEEEEEEEEEEDEESGNQSDRSGSSGI KWRKDSPWVKPSRKRRKREPPRAKEPR GSSGPSEYMEVPLGSLELPSEGTLSPNHL TSSLETERGFEELPLCSCRMEAPKIDRISI CMATESVDGELSGCNAAILKRETMRPSS VLCETHRARMVKHHCCPGCGYFCTAG	
SRSRSHSRGHRRASRDRSAKYKFSRERA ESGRSERGPPDWRLESSNGKMASRRSEI DLLNRMIVWKHGLLI 3372 A 239 3348 PMQNCMCSLTLSVLPLGPQPPVPEKRPP MSDDVHSLGKVTSDLAKRRKLTS*GGI ARRSGEVTLTKGDPGSLEEWETVVGDD SYSVDERVDSDSKSEVEALTEQLSEEEE EEEEEEEEEEEEEEEEEEEEEEEEEEEEEEE	
ESGRSERGPPDWRLESSNGKMASRRSEI DLLNRMIVWKHGLLI 3372 A 239 3348 PMQNCMCSLTLSVLPLGPQPPVPEKRPP MSDDVHSLGKVTSDLAKRRKLTS*GGI ARRSGEVTLTKGDPGSLEEWETVVGDD SYSVDERVDSDSKSEVEALTEQLSEEEE EEEEEEEEEEEEEDEESGNQSDRSGSSGI KWRKDSPWVKPSRKRRKREPPRAKEPR GSSGPSEYMEVPLGSLELPSEGTLSPNHL TSSLETERGFEELPLCSCRMEAPKIDRISI CMATESVDGELSGCNAAILKRETMRPSS VLCETHRARMVKHHCCPGCGYFCTAG	
DLLNRMIVWKHGLLI 3372 A 239 3348 PMQNCMCSLTLSVLPLGPQPPVPEKRPP MSDDVHSLGKVTSDLAKRRKLTS*GGI ARRSGEVTLTKGDPGSLEEWETVVGDD SYSVDERVDSDSKSEVEALTEQLSEEEE EEEEEEEEEEEEEEEDEESGNQSDRSGSSGI KWRKDSPWVKPSRKRRKREPPRAKEPR GSSGPSEYMEVPLGSLELPSEGTLSPNHL TSSLETERGFEELPLCSCRMEAPKIDRISI CMATESVDGELSGCNAAILKRETMRPSS VLCETHRARMVKHHCCPGCGYFCTAG	
3372 A 239 3348 PMQNCMCSLTLSVLPLGPQPPVPEKRPP MSDDVHSLGKVTSDLAKRRKLTS*GGI ARRSGEVTLTKGDPGSLEEWETVVGDD SYSVDERVDSDSKSEVEALTEQLSEEEE EEEEEEEEEEEEEEEDEESGNQSDRSGSSGI KWRKDSPWVKPSRKRRKREPPRAKEPR GSSGPSEYMEVPLGSLELPSEGTLSPNHL TSSLETERGFEELPLCSCRMEAPKIDRISI CMATESVDGELSGCNAAILKRETMRPSS VLCETHRARMVKHHCCPGCGYFCTAG	
MSDDVHSLGKVTSDLAKRKLTS*GGI ARRSGEVTLTKGDPGSLEEWETVVGDD SYSVDERVDSDSKSEVEALTEQLSEEEE EEEEEEEEEEEEEEEDEESGNQSDRSGSSGI KWRKDSPWVKPSRKRRKREPPRAKEPR GSSGPSEYMEVPLGSLELPSEGTLSPNHL TSSLETERGFEELPLCSCRMEAPKIDRISI CMATESVDGELSGCNAAILKRETMRPSS VLCETHRARMVKHHCCPGCGYFCTAG	EIOHFR
ARRSGEVTLTKGDPGSLEEWETVVGDD SYSVDERVDSDSKSEVEALTEQLSEEEE EEEEEEEEEEEEEEEEGNQSDRSGSSGF KWRKDSPWVKPSRKRRKREPPRAKEPR GSSGPSEYMEVPLGSLELPSEGTLSPNHL TSSLETERGFEELPLCSCRMEAPKIDRIST CMATESVDGELSGCNAAILKRETMRPSS VLCETHRARMVKHHCCPGCGYFCTAG	
SYSVDERVDSDSKSEVEALTEQLSEEEE EEEEEEEEEEEEEEEEEEEGNQSDRSGSSGF KWRKDSPWVKPSRKRRKREPPRAKEPR GSSGPSEYMEVPLGSLELPSEGTLSPNHL TSSLETERGFEELPLCSCRMEAPKIDRIST CMATESVDGELSGCNAAILKRETMRPSS VLCETHRARMVKHHCCPGCGYFCTAG	
EEEEEEEEEEEEEDEESGNQSDRSGSSGI KWRKDSPWVKPSRKRRKREPPRAKEPR GSSGPSEYMEVPLGSLELPSEGTLSPNHL TSSLETERGFEELPLCSCRMEAPKIDRISI CMATESVDGELSGCNAAILKRETMRPSS VLCETHRARMVKHHCCPGCGYFCTAG	
KWRKDSPWVKPSRKRRKREPPRAKEPR GSSGPSEYMEVPLGSLELPSEGTLSPNHA TSSLETERGFEELPLCSCRMEAPKIDRISI CMATESVDGELSGCNAAILKRETMRPS VLCETHRARMVKHHCCPGCGYFCTAG	
GSSGPSEYMEVPLGSLELPSEGTLSPNHA TSSLETERGFEELPLCSCRMEAPKIDRISI CMATESVDGELSGCNAAILKRETMRPSS VLCETHRARMVKHHCCPGCGYFCTAG	
TSSLETERGFEELPLCSCRMEAPKIDRISI CMATESVDGELSGCNAAILKRETMRPS VLCETHRARMVKHHCCPGCGYFCTAG	
CMATESVDGELSGCNAAILKRETMRPSS VLCETHRARMVKHHCCPGCGYFCTAG	
VLCETHRARMVKHHCCPGCGYFCTAG	
DIKANIKTIKACABODINGWATCHICON	
QEVTIPRGDGVTPPAGTAAPAPPPLSQD	
TSQPSARMRGHGEPRRPPCDPLADTIDS	
PNGGCLSAVGLPLGPGREALEKALVIQE	
LRFHPRQLYLSVKQGELQKVILMLLDNI	
DQQSKRTPLHAAAQKGSVEICHVLLQA	GANINIA
VDKQQRTPLMEAVVNNHLEVARYMVQ	
YSKEEDGSTCLHHAAKIGNLEMVSLLLS	
VNAQDSGGWTPIIWAAEHKHIEVIRMLI	
VTLTDNEENICLHWASFTGSAAIAEVLL	
HAVNYHGDTPLHIAARESYHDCVLLFL	
ELRNKEGDTAWDLTPERSDVWFALQL	
GVGNRAIRTEKIICRDVARGYENVPIPCV	
EPCPEDYKYISENCETSTMNIDRNITHLO	
DDCSSSNCLCGQLSIRCWYDKDGRLLQ	•
PLIFECNQACSCWRNCKNRVVQSGIKVI	
TAKMGWGVRALQTIPQGTFICEYVGEL	
VREDDSYLFDLDNKDGEVYCIDARYYG	
HLCDPNIPVRVFMLHQDLRFPRIAFFSS	
ELGFDYGDRFWDIKSKYFTCQCGSEKC	
ALEQSRLARLDPHPELLPELGSLPPVNT	
<u> </u>	
	KHSAEAI
FANFVDFNPSGTCIASAGSDQTVKVWD	KHSAEAI FSDSVG
LQHYQVHSGGVNCISFHPSGNYLITASSI DV L KODL NUTL OCHTODVETVOESKOCE	KHSAEAI FSDSVG VRVNKL
DLLKGRLIYTLQGHTGPVFTVSFSKGGE	KHSAEAI FSDSVG VRVNKL DGTLKIL
ADTQVLLWRTNFDELHCKGLTKRNLKF	KHSAEAI FSDSVG VRVNKL DGTLKIL LFASGG
PHLLDIYPRTPHPHEEKVETVEDFFLHLI	KHSAEAI FSDSVG VRVNKL DGTLKIL LFASGG LLHFDSP
R*SICRSLLPLLWISFLLILPQQQKPVVGI	FSDSVG VRVNKL DGTLKIL LFASGG KLHFDSP RLIQSL

SEQ ID NO:	Method	Predicted beginning nucleotide location corresponding to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion
				KRPVDIS*TLP*CHQNVCQQPRKRKQKT*VTSPV KVK/VSIPLAVTDALEHIMEQLNVLTQTVSILEQR LTLTEDKLKDCLENQQKLFSAVQQKS
3374	A	398	21	WLYPMALSILDIKMSPSWYFHMAIGIINWNTTAG LSGTLYPKVPQKYILFDSVILLLGMLRKIRQVCQ NVYMKGCSPITLFKIVHYWPGAVAHAYNPSTLG GQVG/WQIT*GQEFETSLDYMVKPHLY
3375	A .	3	1051	VPTQQILAFPEQTNTKDWTVTPEHVLPESQSLLT FEEVAMYFSQEEWELLDPTQKALYNDVMQENY ETVISLALFVLPKPKVISCLEQGEEPWVQVSPEFK DSAGKSPTGLKLKNDTENHQPVSLSDLEIQASAG VISKKAKVKVPQKTAGKENHFDMHRVGKWHQ DFPVKKRKKLSTWKQELLKLMDRHKKDCAREK PFKCQECGKTFRVSS\DL\IKHQRIHTEEKPYKCQ QCDKRFRWSSDLNKHLTTHQGIKPYKCSWGGKS FSQNTNLHTHQRTHTGEKPFTCHECGKKFSQNS HLIKHRRTHTGEQPYTCSICRRNFSRRSSLLRHQK LHL*REACPVSHFWKTF
3376	A	137	2329	SFESPAPLPSTCFPQERQDPGPCYVSGAMAGLGP GVGDSEGGPRPLFCRKGALRQKVVHEVKSHKFT ARFFKQPTFCSHCTDFIWGIGKQGLQCQVCSFVV HRRCHEFVTFECPGAGKGPQTDDPRNKHKFRLH SYSSPTFCDHCGSLLYGLVHQGMKCSCCEMNVH RRCVRSVPSLCGVDHTERRGRLQLEIRAPTADEI HVTVGEARNLIPMDPNGLSDPYVKLKLIPDPRNL TKQKTRTVKATLNPVWNETFVFNLKPGDVERRL SVEVWDWDRTSRNDFMGAMSFGVSELLKAPVD GWYKLLNQEEGEYYNVPVADADNCSLLQKFEA CNYPLELYERVRMGPSSSPIPSPSPSPTDPKRCFFG ASPGRLHISDFSFLMVLGKGSFGKVMLAERRGSD ELYAIKILKKDVIVQDDDVDCTLVEKRVLALGG RGPGGRPHFLTQLHSTFQTPDRLYFVMEYVTGG DLMYHIQQLGKFKEPHAAFYAAEIAIGLFFLHNQ GIIYRDLKLDNVMLDAEGHIKITDFGMCKENVFP GTTTRTFCGTPDYIAPEIIAYQPYGKSVDWWSFG VLLYEMLAGQPPFDGEDEEELFQAIMEQTVTYP KSLSREAVAICKGFLTKHPGEAPGASGP*WGNLT IRAHGFFPLGFDWERLERL\EIPASFSRPRPCGPQR RGIFDKFFTRAAPA\LTPPARLVLDSIDQADFQGF
3377	A	918	738	SSMLWGFSVFRRSWILNCWLSSSQVGISAACKFS TLTHTHTHTHTRHAPFCGTCLYY
3378	A	1126	456	FSKLIMKTFIIGISGVTNSGKTTLAKNLQKHLPNC SVISQDDFFKPESEIETDKNGFLQYDVLEALNME KMMSAISCWMESARHSVVSTDQESAEEIPILIIEG FLLFNYKPLDTIWNRSYFLTIPYEECKRRRSTRVY QPPDSPGYFDGHVWPMYLKYRQEMQDITWEVV YLDGTKSEEDLFLQVYEDLIQELAKQKCLQVTA* RRNTTNPS/CK*IRKLQGVI
3379	A	1126	456	FSKLIMKTFIIGISGVTNSGKTTLAKNLQKHLPNC SVISQDDFFKPESEIETDKNGFLQYDVLEALNME KMMSAISCWMESARHSVVSTDQESAEEIPILIIEG FLLFNYKPLDTIWNRSYFLTIPYEECKRRRSTRVY QPPDSPGYFDGHVWPMYLKYRQEMQDITWEVV YLDGTKSEEDLFLQVYEDLIQELAKQKCLQVTA*

SEQ ID NO:	Method	Predicted beginning nucleotide location corresponding to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion
.3380	A	1443	794	RRNTTNPS/CK*IRKLQGVI ARRGELAGGGRASGGRSGGDGGGGGGARAPEG VRAPAAGQPRATKGAPPPPGTPPPSPMSSAIERKS LDPSEEPVDEVLQIPPSLLTCGGCQQNIGDRYFLK AIDQYWHEDCLSCDLCGCRLGEVGRRLYYKLGR KLCRRDYLRLFGQDGLCASCDKRIRAYEMTMRV KDKVYHLECFKCAACQKHFCVGDRYLLINSDIV CEQDIYEWTKINGMI
3381	A .	945	474	SLKLRKPPLPTDGVHFVFVESQLDFWGPQEMLT QQGMALQNYDNKLVKCIEELCQKQEELCWQIQ QEEDKKQRLQNEVRQLTEKLACVNEKLARVNE NLARKIASCSKFYQTIAETEATYLKILESF*\TLLS VRKREAGNLTKATAPDQKSSGGRDS
3382	A		1458	GIRGKMADRGGVGEAAAVGASPASVPGLNPTLG WRERLRAGLAGTGASLWFVAGLGLLYALRIPLR LCENLAAVTVFLNSLTPKFYVALTGTSSLISGLIFI FEWWYFHKHGTSFIEQVSVSHLQPLMGGTESSIS EPGSPSRNRENETSRQNLSECKVWRNPLNLFRGA EYRRYTWVTGKEPLTYYDMNLSAQDHQTFFTC DTDFLRPSDTVMQKAWRERNPPARIKAAYQALE LN/E*LCHCICSTG*GRSNNYCRC*KVI*TGTQGR RNNL*AVTAVPAPKSSA*SSTEERYQCTGIY*LKI GNVCKKIRKNKRSSKNNERFDE*ISSSYHVEHP* KSL\KSLLELQAYPDVQAVLAKYDDISLPKSAAIC YTAALLKTRTVSEKFSPETASTRGLSAAEINAVD AIHRAVEFNPHVPKYLLEMKSLILPPEHILKRGDS EAIAYAFFHLQHWKRIEGALNLLQCTWEGSKYS FPKVTLISLTIH
3383	A	282	2443	RGKGFKEFFLGVCQTFIPCLCAEGIQLQFFCSGSG SSPLLKDLESMKTGLFFLCLLGTAAAIPTNARLLS DHSKPTAETVAPDNTAIPSLRAEAEENEKETAVS TEDDSHHKAEKSSVLKSKEESHEQSAEQGKSS\S QELGIEGFKRDSDGSL*VWNL\EYGTNLKGTLDI KEDMSEPQEKKLSENTDFLAPGVSSFTDSNQQES ITKREENQEQPRNYSHHQLNRSSKHSQGLRDQG NQEQDPNISNGEEEEEKEPGEVGTHNDNQERKTE \LPREHANSKQEEDNTQSDDILEESDQPTQVSKM QEDEFDQGNQEQEDNSNAEMEEENASNVNKHIQ ETEWQSQEGKTGLEAISNHKETEEKTVSEALLME PTDDGNTTPRNHGVDDDGDDDGDDGGTDGPRH SA\SDDYFHPKPGLFWEAERA\HSIAYSPSKLREQ REKVHENENIGTTEPGEHQEAKKAENSSNEEETS SEGNMR\VHAVDSCMSFQCKRGHICKADQQGKT SLVSCQDPVT\CPPTKPLDQVCGTDNQTYASSCH LFATKCRLEGTKKGHQLQLDYFG\ASKSIPT\CRD FEVIQ\FPLRMRDW\LKNILMQLYEANSEHAGYL NEK\QRNKVKKIYL\DEKRLLAGDHPIDLLLRDFK KNYHMYVYPVHWQFSELDQHPMDRVLTHSELA PLRASLVPMEHCITRFFEECDPNKDKHITLKEWG HCFGIKEEDIDENLLF
3384	A	3166	928	PSRPHPTHAAMAGPEGFQYRALYPFRRERPEDLE LLPGDVLVVSRAALQALGVAEGGERCPQSVGW MPGLNERTRQRGDFPGTYVEFLGPVALARPGPR PRGPRPLPARPRDGAPEPGLTLPDLPEQFSPPDVA PPLLVKLVEAIERTGLDSESHYRPELPAPRTDWSL

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SEQ ID Me NO:	ethod	Predicted beginning nucleotide location corresponding to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion
				SDVDQWDTAALADGIKSFLLALPAPLVTPEASAE ARRALREAAGPVGPALEPPTLPLHRALTLRFLLQ HLGRVASRAPALGPAVRALGATFGPLLLRAPPPP SSPPPGGAPDGSEPSPDFPALLVEKLLQEHLEEQE VAPPALPPKPPKAK\PASTVPGPNGGSPPSL\QDA EWYWGD\ISREEVNEKLRDTPDGTFLVRDASSKI QGEYTLTLRKGGNNKLIKVFHRDGHYGFSEPLTF CSVVDLINHYRHESLAQYNAKLDTRLLYPVSKY QQDQIVKEDSVEAVGAQLKVYHQQYQDKSREY DQLYEEYTRTSQELQMKRTAIEAFNETIKIFEEQG QTQEKCSKEYLERFRREGN\QTKEMQRILLNSER LKSRIA\EIHESRT\KL\EQQLLVPRASDNKRD\IDK PH*TSLKPDLMQLRKIRDQYLVWLTQKGARQKK INEWLGIKNETEDQYALMEDEDDLPHHEERTWY VGKINRTQAEEMLSGKRDGTFLIRESSQRGCYAC SVVVDGDTKHCVIYRTATGFGFAEPYNLYGSLK ELVLHYQHASLVQHNDALTVTLAHPVRAPGPGP
	j			PPAAR
3385 A		43	2372	TRDVNSWKELCFNHYNKETTNCYRTTRKWTNY KIJEGPFRELRSQGNQVILNLGKERCQLRETGLK LYLPGMDSARHHISHSTSAGPIPSQKEEEMTESQ GTVTFKDVAIDFTQEEWKRLDPAQRKLYRNVML *NYNNLITVGYPFTKPDVIFKLEQEEKPWVMEEE VLRHWQGEIWGVDEHQKNQDRLLRQVEVKFQ KTLTEEKGNECQKKFANVFPLNSDFFPSRHNLYE YDLFGKCLEHNFDCHNNVKCLMRKEHCEYNEP VKSYGNSSSHFVITPFKCNHCGKGFNQTLDLIRH LRIHTGEKPYECSNCRKAFSHKEKLIKHYKIHSRE QSYKCNECGKAFIKMSNLIRHQRIHTGEKPYACK ECEKSFSQKSNLIDHEKIHTGEKPYECNECGKAFS QKQSLIAHQKVHTGEKPYACNECGKAFPRIASLA LHMRSHTGEKPYKCDKCGKAFSQFSMLIIHVRIH TGEKPYECNECGKAFSQSSALTVHMRSHTGEKP YECKECRKAFSHKKNFITHQKIHTREKPYECNEC GKAFIQMSNLVRHQRIHTGEKPYICKECGKAFSQ KSNLIAHEKIHSGEKPYECNECGKAFSQKONFIT HQKVHTGEKPYDCNECGKAFSQIASLTLHLRSHT GEKPYECDKCGKAFSQCSLLNLHMRSHTGEKPY VCNECGKAFSQRTFLIVHMRGHTGEKPYECNEC GKAFIQKSTLIHIRGHTGEKPYECKECRKAFSHK KNFITHQKIHTRE/KPFKCNHCGKGFNQTLDLIRH LRIHTGEKPYECSNCRKAFSHKEKLIKHYKIHSRE QSYKCNECGKAFIKMSNLIRHQRIHTGEKPYACK ECEKSFSQKSNLIDHEKIHTGEKPYECNECGKAFS QKQSLIAHQKVHTGEKPYACNECGKAFPRIASLA LHMRSHTGEKPYKCDKCGKAFSQFSMLIIHVRIH TGEKPYECNECGKAFSQSSALTVHMRSHTGEKP YECKECRKAFSHKKNFITHQKIHTREKPYECNEC GKAFIQMSNLVRHQRIHTGEKPYICKECGKAFSQ KSNLIAHEKIHSGEKPYECNECGKAFSQFSMLIIHVRIH TGEKPYECNECGKAFSQSSALTVHMRSHTGEKP YECKECRKAFSHKKNFITHQKIHTREKPYECNEC GKAFIQMSNLVRHQRIHTGEKPYICKECGKAFSQ KSNLIAHEKIHSGEKPYECNECGKAFSQKONFIT HQKVHTGEKPYDCNECGKAFSQIASLTLHLRSHT GEKPYECDKCGKAFSQCSLLNLHMRSHTGEKPY VCNECGKAFSQRTFLIVHMRGHTGEKPYECNEC GKAFSQSSSLTIHIRGHTGEKPYECKECRKAFSHK KNFITHQKIHTRENPLSVIIVEKASIRLWTSSDI

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SEQ ID NO:	Method	Predicted beginning nucleotide location corresponding to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion
3386	A	201	1032	WDDYPQGALRRREAAEGLHFLGPPGRVRGQLR GITGPAWYCHSPSHSLLSAFCHLPTPSRCPAMAR PPVPGSVVVPNWHES/RRGQGVPGLHSAQEPPAG VWAA*AASAAAA\LSIDTASYKIFVSGKSGVGKT ALVAKLAGLEVPVVHHETTGIQTTVVFWPAKLQ ASSRVVMFRFEFWDCGESALKKFDHMLLACME NTDAFLFLFSFTDRASFEDLPGQLARIAGEAPGV VRMVIGSKFDQYMHTDVPERDLTAFRQAWELPL LRVKSVPGRRLG
3387	A	86	96	GSSPDPASLITMKNQDKKNGAAKQSNPKSSPGQP EAGPEGAQERPSQAAPAVEAEGPGSSQAPRKPEG AQARTAQSGALRDVSEELSRQLEDILSTYCVDNN QGGPGEDGAQGEPAEPEDAEKSRTYVARNGEPE PTPVVNGEKEPSKGDPNTEEIRQSDEVGDRDHRR PQEKKKAKGLGKEITLLMQTLNTLSTPEEKLAAL CKKYAELLEEHRNSQKQMKLLQKKQSQLVQEK DHLRGEHSKAVLARSKLESLCRELQRHNRSLKE EGVQRAREEEEKRKEVTSHFQVTLNDIQLQMEQ HNERNSKLRQENMELAERLKKLIEQYELREEHID KVFKHKDLQQQLVDAKLQQAQEMLKEAEERHQ REKDFLLKEAVESQRMCELMKQQETHLKQQLA LYTEKFEEFQNTLSKSSEVFTTFKQEMEKMTKKI KKLEKETTMYRSRWESSNKALLEMAEEKTVRD KELEGLQVKIQRLEKLCRALQT/GAQ*PVRGQRW GSHRTSAVRIFS
	A	98	3197	ARPEVPAPPA WLSRRGAAKMGDKKDDKDSPKK NKGKERRDLDDLKKEVAMTEHKMSVEEVCRKY NTDCVQGLTHSKAQEILARDGPNALTPPTTPEW VKFCRQLFGGFSILLWIGAILCFLAYGIQAGTEDD PSGDNLYLGIVLAAVVIITGCFSYYQEAKSSKIME SFKNMVPQQALVIREGEKMQVNAEEVVVGDLV EIKGGDRVPADLRIISAHGCKVDNSSLTGESEPQT RSPDCTHE\NPLKTRNITFFSNNFVEGTARGVVVA TGDRTVMGRIATLASGLEVGKTPIAIEIEHFIQLIT GVAVFLGVSFFILSLILGYTWLEAVIFLIGIIVANV PEGLLATVTVCLTLTAKRMARKNCLVKNLEAVE TLGSTSTICSDKTGTLTQNRMTVAHMWFDNQIH EADTTEDQSGTSFDKSSHTWVALF*H/LLGFCNR PVFKGGQDNIPVLKRDVAGDASESALLKCIELSS GSVKLMRERNKKVAEIPFNSTNKYQLSIHETEDP NDNRYLLVMKGAPERILDRCSTILLQGKEQPLDE EMKEAFQNAYLELGGLGERVLGFCHYYLPEEQF PKGFAFDCDDVNFTTDNLCFVGLMSMIGPPRAA VPDAVGKCRSAGIKVIMVTGDHPITAKAIAKGV GIIFEGNETVEDIAARLNIPVSQVNPRDAKACVIH GTDLKDFTSEQIDEILQNHTEIVFARTSPQQKLIIV EGCQRQGAIVAVTGDGVNDSPALKKADIGVAM GIAGSDVSKQAADMILLDDNFASIVTGVEEGRLI FDNLKKSIAYTLTSNIPEITPFLLFIMANIPLPLGTI TILCIDLGTDMVPAISLAYEAAESDIMKRQPRNPR TDKLVNERLISMAYGQIGMIQALGGFFSYFVILA ENGFLPGNLVGIRLNWDDRTVNDLEDSYGQQW TYEQRKVVEFTCHTAFFVSIVVVQWADLIICKTR RNSVFQQGMKNKILIFGLFEETALAAFLSYCPGM DVALRMYPLKPSWWFCAFPYSFLIFVYDEIRKLI

Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, Predicted end SEQ ID Method Predicted E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, nucleotide NO: beginning nucleotide location corresponding N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, location to last amino T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, corresponding acid residue of X=Unknown, *=Stop codon, /=possible nucleotide deletion, to first amino peptide acid residue of \=possible nucleotide insertion peptide sequence sequence LRRNPGGWVEKETYY 5250 VERLLGCRNSKRTWRMLISKNMPWRRLQGISFG 45 3389 Α MYSAEELKKLSVKSITNPRYLDSLGNPSANGLYD LALGPADSKEVCSTCVQDFSNCSGHLGHIELPLT VYNPLLFDKLYLLLRGSCLNCHMLTCPRAVIHLL LCQLRVLEVGALQAVYELERILNRFLEENPDPSA SEIREELEQYTTEIVQNNLLGSQGAHVKNVCESK SKLIALFWKAHMNAKRCPHCKTGRSVVRKEHNS KLTITFPAMVHRTAGQKDSEPLGIEEAQIGKRGY LTPTSAREHLSALWKNEGFFLNYLFSGMDDDGM ESRFNPSVFFLDFLVVPPSRYRPVSRLGDQMFTN GOTVNLQAVMKDVVLIRKLLALMAQEQKLPEE VATPTTDEEKDSLIAIDRSFLSTLPGQSLIDKLYNI WIRLQSHVNIVFDSEMDKLMMDKYPGIRQILEK KEGLFRKHMMGKRVDYAARSVICPDMYINTNEI GIPMVFATKLTYPQPVTPWNVQELRQAVINGPN VHPGASMVINEDGSRTALSAVDMTQREAVAKQ LLTPATGAPKPQGTKIVCRHVKNGDILLLNRQPT LHRPSIOAHRARILPEEKVLRLHYANCKAYNADF DGDEMNAHFPOSELGRAEAYVLACTDQQYLVP KDGQPLAGLIQDHMVSGASMTTRGCFFTREHYM ELVYRGLTDKVGRVKLLSPSILKPFPLWTGKQVV STLLINIIPEDHIPLNLSGKAKITGKAWVKETPRSV **PGFNPDSMCESQVIIREGELLCGVLDKAHYGSSA** YGLVHCCYEIYGGETSGKVLTCLARLFTAYLQL YRGFTLGVEDILVKPKADVKRQRIEESTHCGPQ AVRAALNLPEAASYDEVRGKWQDAHLGKDQRD FNMIDLKFKEEVNHYSNEINKACMPFGLHRQFPE NTLQLMVQSGAKGSTVNTMQISCLLGQIELEGRS TPLMASGKSLPCFEPYEFTPRAGGFVTGRFLTGIK PPEFFFHCMAGREGLVDTAVKTSRSGYLQRCIIK HLEGLVVQYDLTVRDSDGSVVQFLYGEDGLDIP KTQFLQPKQFPFLASNYEVIMKSQHLHEVLSRAD PKKALHHFRAIKKWQSKHPNTLLRRGAFLSYSQ KIQEAVKALKLESENRNGR/RPWDS/G/RMLRMW YELDEESRRKYOKKAAACPDPSLSVWRPDIYFAS VSETFETKVDDYSQEWAAQTEKSYEKSELSLDR LRTLLQL\KWQRSLCEPGEAVGLLAAQSIGEPST **QMTLNTFHFAGRGEMNVTLGIPRLREILMVASA** NIKTPMMSVPVLNTKKALKRVKSLKKQLTRVCL GEVLQKIDVQESFCMEEKQNKFQVYQLRFQFLP HAYYOQEKCLRPEDILRFMETRFFKLLMESIKKK NNKASAFRNVNTRRATQRDLDNAGELGRSRGE OEGDEEEEGHIVDAEAEEGDADASDAKRKEKQE **EEVDYESEEEEEREGEENDDEDMQEERNPHREG** ARKTQEQDEEVGL/GH*GGPVPSRPPDAAPETHP OPGAPGA\EAMERRVOAVREIHPFIDDYQYDTEE SLWCQVTVKLPLMKINFDMSSLVVSLAHGAVIY ATKGITRCLLNETTNNKNEKELVLNTEGINLPELF KYAEVLDLRRLYSNDIHAIANTYGIEAALRVIEK **EIKDVFAVYGIAVDPRHLSLVADYMCFEGVYKP** LNRFGIRSNSSPLQQMTFETSFQFLKQATMLGSH DELRSPSACLVVGKVVRGGTGLFELKQPLR 2080 ILPPLEGPPAQASPSSTMLGEGSQPDWPGGSRYD 3390 A 2 LDEIDAYWLELINSELKEMERPELDELTLERVLE

SEQ ID NO:	Method	Predicted beginning nucleotide location corresponding to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion
				ELETLCHQNMARAIETQEGLGIEYDEDVVCDVC RSPEGEDGNEMVFCDKCNVCVHQACYGILKVPT GSWLCRTCALGVQPKCLLCPKRGGALKPTRSGT KWVHVSCALWIPEVSIGCPEKMEPITKISHIPASR WALSCSLCKECTGTCIQCSMPSC\VTAFHVTCAF DHGLEMRTILADNDEVKFKSFCQEHSDGGPRNE PTSEPTEPSQAGEDLEKVTLRKQRLQQLEEDFYE LVEPAEVAERLDLAEALVDFIYQYWKLKRKANA NQPLLTPKTDEVDNLAQQEQDVLYRRLKLFTHL RQDLERVRNLCYMVTRRERTKHAICKLQEQIFH LQMKLIEQDLCRAGLSTSFPIDGTFFNSWLAQSV QITAENMAMSEWPLNNGHREDPAPGLLSEELLQ DEETLLSFMRDPSLRPGDPARKARGRTRLPAKK KPPPPPPQDGPGSRTTPDKAPKKTWGQDAGSGK GGQGPPTRKPPRRTSSHLPSSPAAGDCPILATPES PPPLAPETPDEAASVAADSDVQVP\GPAASPKPLG RLRPPPREPR*T\RRLPGC/ARPDAGDGDHLSAVA ERPKV\SLHFDTETDG\YFS\DGEMSNS\DV\EAED GGVQRGPREAGAKE\VVRMGVLAS
3391	A	1555	327	NSFLHFLHLKVRTMFLFPSFPVLLLSVVTASCSKT KACADTQKTCSMITCGIPVTNGTPGRDGRDRPK GEKGEPGLGQVSVAS*ISTSGRCSSKSVLEPATRG LKHRLGEAPLSSGPMLHSEQPL*NAIASKTKLFV DSLGSHISTQELGVCGCPFRGVSCLVGELALVQA LH*VAGESFFFGSDHWLIGCAGGEQEWSIELLGK KKRVTATGSSSLCLATGQGLRGLQGPPGKMGPP GNTGTSGIPGPRGQKGDRGDNSVAEAKLANLER KL*SLRSELDHTKKL*PFSLGK\MSGKKLFVTNGE RMPFSKVKALCAGLQATVAAPKNAEENKAIQDV AKDTAFLGITDEATEGQFMYLTGGRLTYSNWKK DEPNDHGSGEDCVILLNNGLWNGISCTSSFIAICE FPA
3392	A	218	1773	GGSRRNQRRSIPVLGYFLKQKKMTKAQESLTLE DVAVDFTWEEWQFLSPAQKDLYRDVMLENYSN LVSVGYQAGKPDALTKLEQGEPLWTLEDEIHSP AHPEIEKADDHLQQPLQNQKILKRTGQRYEHGR TLKSYLGLTNQSRRYNRKEPAEFNGDGAFLHDN HEQMPTEIEFPESRKPISTKSQFLKHQQTHNIEKA HECTDCGKAFLKKSQLTEHKRIHTGKKPHVCSL CGKAFYKKYRLTEHERAHRGEKPHGCSLCGKAF YKRYRLTEHERAHKGEKPYGCSECGKAFPRKSE LTEHQRIHTGIKPHQCSECGRAFSRKSLLVVHQR THTGEKPHTCSECGKGFIQKGNLNIHQRTHTGEK PYGCIDCGKAFSQKSCLVAHQRYHTGKTPFVCPE CGQPCSQKSGLIRHQKIHSGEKPYKCSDCGKAFL TKTMLIVHHRTHTGERPYGCDECEKAYFYMSCL VKHKRIHSREKRGD/CSEGGKSFHSKSQLKS**TC AGEKPC*YGNCGNGGRAV
3393	A	46	1464	ARSLSGAPSGSSRQDGTSLLRTGAGYSSSQSIETL SLPPGPSHLVGDKSQGGRSCQGQITSAASGKTSK SEPNHVIFKKISRDKSVT\IYLGNRDY\IDHV\SQV QPVDGVVLVDPDLVKGKKVYVTLTCAFRYGQE DIDVIGLTFRRDLYFSRVQVYPPVGAASTPTKLQ ESLLKKLGSNTYPFLLTFPDYLPCSVMLQPAPQD SGKSCGVDFEVKAFATDSTDAEEDKIPKKSSVRL

SEQ ID	Method	Predicted	Predicted end nucleotide	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine,
NO:		beginning nucleotide	location	I=Isoleucine, K=Lysine, L=Leucine, M=Methionine,
	ŀ	location	corresponding	N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine,
		corresponding	to last amino	T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine,
		to first amino	acid residue of	X=Unknown, *=Stop codon, /=possible nucleotide deletion,
ļ		acid residue of	peptide sequence	\ possible nucleotide insertion
		peptide sequence	sequence	
 	<u> </u>	1		LIRKVQHAPLEMGPQPRAEAAWQFFMF\DKPLH
	-			LAVSLNKRDLFPMGSPIPVPVSVP\NNTEKPVKKI
]			1	KA\SVEQVANVVLYS\SDY\YVKPVAMEEAQEKV
	+			PPNSTWTKA\LTLL\PWLVNNRERRGIALDGKIKH
		1		EDTNLASSTIIKEGIDRKRSWEILVSYPDQR*SSTV
				SGFLGRASPSQ*SRPT*RSQFRL\MHPQP\EDPA\K
		İ	Ì	ESYQDANLVF\EEFARP*ILKDAGEA*\EGKRDQE
3394	A	211	1591	RPPTMAADQRPKADTLALRQRLISSSCRLFFPEDP
3371	1		1000	VKIVRAQGQYMYDEQGAEYIDCISNVAHVGHCH
				PLVVQAAHEQNQVLNTNSRYLHDNIVDYAQRLS
		ļ		ETLPEQLCVFYFLNSGSEANDLALRLARHYTGH
				QDVVVLDHAYHGHLSSLIDISPYKFRNLDGQKE
	1			WVHVAPLPDTYRGPYREDHP\THVEDGLEKAFS*
			1	KRVVQGRNRQICRRQIAAFFAESLPSVGGQIIPPA
				GYFSQVAEHIRKAGGVFVADEIQVGFGRVGKHF
		ļ		WAFQLQGKDFVPDIVTMGKSIGNGHPVACVAAT
	1			QPVARAFEATGVEYFNTFGGSPVSCAVGLAVLN
				VLEKEQLQDHATSVGSFLMQLLGQQKIKHPIVG
	1	ł	}	DVRGVGLFIGVDLIKDEATRTPATEEAAYLVSRL
				KENYVLLSTDGPGRNILKFKPPMCFSLDNARQV
<u> </u>	1			VAKLDAILTDMEEKVRSCETLRLQP
3395	A	1	1424	FRDGFSLRCGCNAELPGRGGDDAADRAIQRFLR
3393	A	1	1424	TGAAVRYKVMKNWGVIGGIAAALAAGIYVIWG
				PITERKKRRKGLVPGLVNLGNTCFMNSLLQGLSA
1				CPAFIRWLEEFTSQYSRDQKEPPSHQYLSLTLLHL
1				LKALSCQEVTDDEVLHASCLLDVLRMYRWQISS
			1	
	}			FEEQDAHELFHVITSSLEDERDRQPRVTHLFDVH
	1			SLE\HSQK*LPKQITCRTRGSPHPTSNHWKSQHPF
ļ				HGRLTSNMVCKHCEHQSPVRFDTFDSLSLSIPAA TWGHPLTLDHCLHHFISSESVRDVVCDNCTKIEA
ļ	1	ļ	1	
	1			KGTLNGEKVEHQRTTFVKQLKLGKLPQCLCIHL
				QRLSWSSHGTPLKRHEHVQFNEFLMMDIYKYHL
				LGHKPSQHNPKLNKNPGPTLELQDGPGAPTPGL
				NQPGAPKTQIFMNGACSPSLLPTLSAPMPFPLPV VPDYSSSTYLFRLMGSCRPPWETWHSGTLCSFTD
			1.	
-	 	100	107	GPHL
3396	A	109	107	TQEAGLIFFSPPFSLSLSLSLPLSLFLLSHPHSRTPP
				NRTPRRTRIPQRPAVMYSPLCLTQDEFHPFIEALL
				PHVRAFAYTWFNLQARKRKYFKKHEKRMSKEE
				ERAVKDELLSEKPEVKQKWASRLLAKLRKDIRP
		1	1	EYREDFVLTVTGKKPPCCVLSNPDQKGKMRRID
1	}		1	CLRQADKVWRLDLVMVILFKGIPLESTDGERLV
				KSPQCSNPGLCVQPHHIGVSVKELDLYLAYFVH
}				AADSSQSESPSQAK*R*H*GPARKWDIWGFQ\DS
			1	FVT\SGVF\SVT*A*LRVSQTPI\AAG\TGPNFSLSD
				LESSSYYSMSPGAMRRSLPSTSSTSSTKRLKSVED
			÷	EMDSPGEEPFYTGQGRSPGSGSQSSGWHEVEPG
			1	MPSPTTLKKSEKSGFSSPSPSQTSSLG\TAFTQHHR
1				PVITGTQSKFHIATPSIL\HFPRHSPFFQQPGPYFSH
		1	1	PAIRYHPQETLKEFVQLVCPDAGQQAGQPNGSS
		1	1	QGKVHNPFLPTPMLPPPPPPPMARPVPLPVPDTK
				PPTTSTEGGAASPTSPTTRS/PGRTRPQQPFL/SYG
				PP*PSNALIGGGGGGAGERAGERADLEM
3397	A	1	2002	TGTLTEDGLDVMGVVPLKGQAFLPLVPEPRRLP
.				VGPLLRALATCHALSRLQDTPVGDPMDLKMVES
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CEO IN	1-32-0-3	I was a second	I D - 31-4-3 - 3	The state of the s
SEQ ID NO:	Method	Predicted beginning nucleotide location corresponding to first amino acid residue of peptide	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \ -possible nucleotide insertion
	<u> </u>	sequence		
				TGWVLEEPAADSAFGTQVLAVMRPPLWEPQLQ AMEEPPVPVSVLHRFPFSSALQRMSVVVAWPGA TQPEAYVKGSPELVAGLCNPETVPTDFAQMLQS YTAAGYRVVALASKPLPSVPSLEAAQQLTRDTV EGDLSLLGLLVMRNLLKPQTTPVIQALRRTRIRA VMVTGDNLQTAVTVARGCGMVAPQEHLIIVHA THPERGQPASLEFLPMESPTAVNGVKDPDQAAS YTVEPDPRSRHLALSGPTFGIIVKHFPKLLPKVLV QGTVFARMAPEQKTELVCELQKLQYCVGMCGD GANDCGALKAADVGISLSQAEASVVSPFTSSMA SIECVPMVIREGRCSLDTSFSVFKYMALYSLTQFI
				SVLILYTINTNLGDLQFLAIDLVITTTVAVLMSRT GPALVLGRVRPPGALLSVPVLSSLLLQMVLVTG VQLGGYFLTLAQPWFVPLNRTVAAPDNLPNYEN TVVFSLSSFQYLILAAAVSKGAPFR\RPLTNNVPF LLASAL*SSVLVVLVLSPGLLHGPLALRNITDTGF KLLLVGLVTLNFVGGLHAGERARPVPPRLPAPPP AQAG\SKKRFKQLERELAEQPWPPLPAGPLR
3398	A	758	1368	FPFRMLTGYLYLMWRRKAFWSGTQRHPLPGGL KRRRPGRGPWPAPGGQGVGPSAL*KAGSPPAN RPGQGE/PGLISPKPVTEVLPDVQGAPVPVPPLPT PPSLPHLQNQPP/TVQHYLLSFSWKPSQGPE*RA* PSPLPPAAMRPDG*PGPASQGPDQPG\PCPPASLP TSPPGKGFQKTETRKHPPPRQQHKPKCTANRPLA SFL
3399	A	906	1091	HHHHHHHHHHHLVAFGKVQ*LQNSPSSSSS SSGCFWQARFSSYRTLHHHHHHHHHHHHH
3400	A	1838	325	PFLSVHRSPHGPSKLCDDPQASLVPEPVPGGCQE PEEMSWPPSGEIASPPELPSSPPPGLPEVAPDATST GLPDTPAAPETSTNYPVECTEGSAGPQSLPLPILE PVKNPCSVKDQTPLQLSVEDTTSPNTKPCPPTPTT PETSPPPPPPPPSSTPCSAHLTPSSLFPSSLESSSEQ KFYNFVILHARADEHIALRVSGRSWEALGVPDG ATFCEDFQVPGRGELSCLQDAIDHSAFIILLLT\SN \FDCR\LSLHQVNQAMMSNLT\RQGSQDCVIP\FLP \LESSPARLSSDTASLLSGLVRLDEHSQIFARKVA NTFKPHRLQARKAMWRKEQDTRALREQSQHLD GERMQAAALNAAYSAYLQSYLSYQAQMEQLQV AFGSHMSFGTGAPYGARMPFGGQVPLGAPPPFP TWPGCPQPPPLHAWQAGTPPPPSPQPAAFPQSLP FPAVPKPFPTASTAPPSEPKGWQP\LIIHHAQMVT SWG*NKH\MWNQRGSQAPEDKTQEAE
3401	A	153	1389	EWGWLGAAQPPEEEAEAEDQESPSSLCREALAEI KKEISPLFIGMEKCSVGGLELTEQTPALLGNMAM ATSLMDIGDSFGHPACPLVSRSRNSPVEDDDDDD DVVFIESIQPPSISAPAIADQRNFIFASSKNEKPQG NYSVIPPSSRDLASQKGNISETIVIDDEEDIETNGG AEKKSSCFIEWGLPGTKNKTNDLDFSTSSLSRSK VNAGMGNSGITTELTLKYIITNVTTLETGISSVNA GQDVNIIITYKTSL*NTNLGDVAKGLQSSNFGVNI QTYTPSLTPQTKTGVNLLTLVE*MWQETYFRME NLQLII/CPEDASTKKANVILPVESSKSFQEFYSTS CLSPCENNWNLKKGVFNKSRCTICSKLAEVWIFI
2400		1.50	1000	PKLLFRLTVIILTFKCYYVLFHLHNARVLDV
3402	A	153	1389	EWGWLGAAQPPEEEAEAEDQESPSSLCREALAEI

SEQ ID NO:	Method	Predicted beginning nucleotide location corresponding	Predicted end nucleotide location corresponding to last amino	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine,
		to first amino acid residue of peptide sequence	acid residue of peptide sequence	X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion
				KKEISPLFIGMEKCSVGGLELTEQTPALLGNMAM ATSLMDIGDSFGHPACPLVSRSRNSPVEDDDDDD DVVFIESIQPPSISAPAIADQRNFIFASSKNEKPQG
				NYSVIPPSSRDLASQKGNISETIVIDDEEDIETNGG AEKKSSCFIEWGLPGTKNKTNDLDFSTSSLSRSK VNAGMGNSGITTELTLKYIITNVTTLETGISSVNA
				GQDVNIIITYKTSL*NTNLGDVAKGLQSSNFGVNI QTYTPSLTPQTKTGV\NLLTLVE*MWQETYFRME NLQLII/CPEDASTKKANVILPVESSKSFQEFYSTS
				CLSPCENNWNLKKGVFNKSRCTICSKLAEVWIFI PKLLFRLTVIILTFKCYYVLFHLHNARVLDV
3403	A	609	2765	SRHCTPAERQNETHRAPDFAMSAVLGHQPPFFPA LTLPPNGAAALSLPGALAKPIMDQLVGAAETGIP FSSLGPQAHLRPLKTMEPEEEVEDDPKVHLEAKE
				LWDQFHKRGTEMVITKSGRRMFPPFKVRCSGLD KKAKYILLMDIIAADDCRYKFHNSRWMVAGKA
				DPEMPKRMYIHPDSPATGEQWMSKVVTFHKLKL TNNISDKHGFTILNSMHKYQPRFHIVRANDILKLP YSTFRTYLFPETEFIAVTAYQNDKITQLKIDNNPF
				AKGFRDTGNGRREKRKQLTLQSMRVFDERHKK ENGTSDESSSEQAAFNCFA\QASSPAA\PL*RTSNL
				KDF\SPSRG*RATPEAEEQRGSTAPRPATRAKISP HPRRRSPAVTRAAPAVKAHLFAAERPRDSGRLD KASPDSRHSPATISSSTRGLGAEERRSPVREG\QA
		*		PAKVEEARALPGKEAFAPLTVQTDAAAAHLAQG PLPGLGFAPGLAGQQFFNGHPLFLHPSQFAMGG AFSSMAAAGMGPLLATVSGASTGVSGLDSTAM
				ASAAAAQGLSGASAATLPFHLQQHVLASQGLA MSPFGSLFPYPYTYMAAAAAA/SSAAASASVHRT
				P\FNLNTMRPRLRYSPYSIPVPVPDGSSLLTTALPS MAAAAGPLDGKAAALAASPAS\VAVDSGSELNS RSS\TLSSSSMSLSPKLCAEKEAATSELQSIQRLVS
3404	A	1082	1308	GLEAKPDRSRSASP LKKFLEVPQSYSLLLSSPFLQ\WRA*RPQNAIG*Q
		1550		FIIKTLVFFGIMRSAGDVLSTQVSCALRIMRTAGC SHSSP
3405	A	1553	559	PRPPTQRLSRFAPPCRTAEFPFRRRAVVTRPAPPR ACTVVGRSSPVTGLAVGAAVAMLTVAARSRPFA PVLSATSRGVAGALT\P*MQATVPATPEQPVLDL
				KRPFLSRESLSGQAVRRPLVASVGLNVPASVCYS HTDIKVPDFSEYRRLEVLDSTKSSRESSEARKGFS YLVTGVTTVGVAYAAKNAVTQFVSSMSASADV
				LALAKIEIKLSDIPEGKNMAFKWRGKPLFVRHRT QKEIEQEAAVELSQLRDPQHDLDRVKKPEWVILI
				GVCTHLGCVPIANAGDFGGYYCPCHGSHYDASG RIRLGPAPLNLEVPTYEFTSDDMVIVG
3406	A	83	2671	CLYPDFCRSVTCAMPCFTHRSCREDPGTSESREM DPVAFKDVAVNFTQEEWALLDISQKNLYREVML ETFWNLTSIGKKWKDQNIEYEYQNPRRNFRSVT
				EEKVNEIKEDSHCGETFTPVPDDRLNFQKKKASP EVKSCDSFVCEVGLGNSSSNMNIRGDTGHKACE
				CQEYGPKPWKSQQPKKAFRYHPSLRTQERDHTG KKPYACKECGKNIIYHSSIQRHMVVHSGDGPYK CKFCGKAFHWLSLYLIHERTHTGEKPYECKQCG KSFSYSATHRIHERTHIGEKPYECQECGKAFHSPR

Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, SEQ ID Method Predicted Predicted end aucleotide E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, NO: beginning I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, nucleotide location corresponding N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, location T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, corresponding to last amino X=Unknown, *=Stop codon, /=possible nucleotide deletion, acid residue of to first amino peptide \=possible nucleotide insertion acid residue of peptide sequence sequence SCHRHERSHMGEKAYQCKECGKAFMCPRYVRR **HERTHSRKKLYECKQCGKALSSLTSFQTHIRMHS** GERPYECKTCGKGFYSAKSFQRHEKTHSGEKPY KCKQCGKAFTRSGSFRYHERTHTGEKPYECKQC GKAFRSAPNLQSHGRTHTGEKPYECKECGKAFIF VNNLQSHERTQTHIRIHSGERRYKCKICGKGFYC PKSFQRHEKTHTGEKLYEC/TATFSSSFSSSSSF*Y HERTHTGEKPYKCEQCGKAFRAVSIL*MHGRTH PEEKPYECEQ*RKAFRSAPHL*IRGRTHNGEKPY ACKKCGKPFGSAQNLRIHERTQTHIMHSVERPYK CKICGRGFYSAKSFQTHEKSYTGEKPYECKQCG KAFVSFTSFRYHERTHTGENPYECKQFGKAFRSV KNLRFHKRTHTGEKPCEYMKRLTLEGNTMNAS NVAKLSLLPVLFNIMKEFTLGRNPISVSNVRKPLF LPLLFNIMKGLTWERNPMSVCHVGKPSFLLVPFN **IMKGLTLERSPMNISNVGKPSDQPRTFKCMEGLT** LEKNPMNVSSMGKRSDLTRFFEYR PAAPSGASPGRVCGVETARPLGVQRRQSADEGP 3407 1426 3 Ā **PGVAGLRHEPPTVWLGSVAHRGTWVCAHRWFG** PAVTRAAQAATMVKLLVAKILCMVGVFFFMLL GSLLPVKIIETDFEKAHRSKKILSLCNTFGGGVFL ATC\LTALLARC*GKSSRRSWSLGHISTDYPL\AE TILLLGFFMTVFLEQLILTFAQENAVLHRPGDLQR RIGRGQRLGV*EPLHGGRAGPRAVRGAPRPRPQP ERAGPLA\PSPVRLLSLAFALSAHSVFEGLALGLQ **EEGEKVVSLFVGVAVHETLVPVALGISMAGSAM** PLRDAAKLAVTVSPMIPLGIGLGLGIEKAQGVPG SVASVLLQGPGGRHLSLFITFPGKSWPRSWRKKS DRLLKVLF\LVVGYTVLAGMGLPQVVSGLAIVPA **AGSPPGAPGRTQAASPGRASPKSEHCGPGPPPVH** KGPPGTRLCPRSYTLSLRALLLFKILLSLKSLYQK 3408 106 4514 EARDRLAQSRAKEKELNSVASELSARQEESEHSH Ā KHLIELRREFKKNVPEEIREMVAPVLKSFQAEVV ALSKRSQEAEAAFLSVYKQLIEAPALWELKLKSR PALGDSRVQQGQHDPKTDNQNTQQKAGFKEGW LAEASEREAFGPGFKDPVPVFEAARSLDDRLQPP SFDPSGQPRRDLHTSWKRNPELLSPKALKATQAE LLELRRKYDEEAASKADEVGLIMTNLEKANQRA EAAQREVESLREQLASVNSSIRLACCSPQGPSGD KVNFTLCSGPRLEAALASKDREILRLLKDVQHLQ SSLQELEEASANQIADLERQLTAKSEAIEKLEEKL QAQSDYEEIKTELSILKAMKLASSTCSLPQGMAK PEDSLLIAKEAFFPTQKFLLEKPSLLASPEEDPSED DSIKDSLGTEQSYPSPQQLPPPPGPEDPLSPSPGQP LLGPSLGPDGTRTFSLSPFPSLASGERLMMPPAAF KGEAGGLLVFPPAFYGAKPPTAPATPAPGPEPLG **GPEPADGGGGGAAGPGAEEEQLDTAEIAFQVKE** QLLKHNIGQRVFGHYVLGLSQGSVSEILARPKP\ WRKLHG**GKEPFIKMKQFLSDEQNVLALRTIQV RORGSITPRIRTPETGSDDAIKSILEQAKKEIESQK GGEPKTSVAPLSIANGTTPASTSEDAIKSILEQAR REMQAQQALLEMEVAPRGRSVPPSPPERPSLAT ASONGAPALVKQEEGSGGPAQAPLPVLSPAAFV **OSIIRKVKSEIGDAGYFDHHWASDRGLLSRPYAS**

Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, Predicted end Method Predicted E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, nucleotide beginning NO: nucleotide location N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, corresponding location T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, corresponding to last amino X=Unknown, *=Stop codon, /=possible nucleotide deletion. acid residue of to first amino \=possible nucleotide insertion acid residue of peptide peptide sequence sequence VSPSLSSSSSSGYSGQPNGRAWPRGDEAPVPPED EAAAGAEDEPPRTGELKAEGATAEAGARLPYYP AYVPRTLKPTVPPLTPEQYELYMYREVDTLELTR OVKEKLAKNGICQRIFGEKVLGLSQGSVSDMLSR PKPWSKLTOKGREPFIRMQLWLSDQLGQAVGQQ **PGASQASPTEPRSSPSPPPSPTEPEKSSQEPLSLSLE** SSKENOOPEGRSSSSLSGKMYSGSQAPGGIQEIV AMSPELDTYSITKRVKEVLTDNNLGQRLFGESIL GLTOGSVSDLLSRPKPWHKLSLKGREPFVRMQL WLNDPHNVEKLRDMKKLEKKAYLKRRYGLIST GSDSESPATRSECPSPCLQPQDLSLLQIKKPRVVL **APEEKEALRKAYQLEPYPSQQTIELLSFQLNLKT** NTVINWFHNYRSRMRREMLVEGTQDEPDLDPSG **GPGILPPGHSHPDPTPQSPDSETEDQKPTVKELEL OEGPEENSTPLTTQDKAQVRIKQEQMEEDAEEE** AGSQPQDSGELDKGQGPPKEEHPDPPGNDGLPK VAPGPLLPGGSTPDCPSLHPQQESEAGERLHPDP LSFKSASESSRCSLEVSLNSPSAASSPGLMMSVSP **VPSSSAPISPSPPGAPPAKVPSASPTADMAGALHP** SAKVNPNLORRHEKMANLNNITYRLERAANREE **ALEWEF** 3409 162 1710 **GPLSPGPYQCRPSLPAQLYPQSLMAAATLRTPTQ GTVTFEDVAVHFSWEEWGLLDEAQRCLYRDVM** LENLALLTSLDVHHQKQHLGEKHFISNVGRALF VKTCTFHVSGEPSTCREVGKDFLAKLGFLHQQA AHTGEQSNSKSDGGAISHRGKTHYNWGEHTKAF SGKHTLVOOORTLTTERCYICSECGKSFSKSYSL NDHWRLHTGEKPYECRECGKSFRQSSSLIQHRR **GHTAVRPHECDECGKLFSNKSNLIKHRRVHTGE** RPYECSECGKSFNQRSALLQHRGVHTGEKPYEC TECGKSFSHNSSLIKHQRIHSG*\RPYECTECGKSF SONSSLIEHHRVHTGERPYKCSECGKSFRQRSAL LOHRGVPTGERPYECSECGKFFPYSSSLGKHQRV HTGSRPYECSECGKSFTQNSGLIKHRRVHTGEKP YECTE*KKSFSHNSSLIKHQRIHSR*KPYE\CKCG N\R*HPGESP*VHSECQ/KSFS*RPYLIECHTVHKG KTLLICRDVQLI 3410 167 789 LCMKGISGGVRVAALAARAEREELPVPAMEPOP Ā TAWGSPHPEAVLQLEVAPESSGPCTDTAKDQQS DKLPDLMPPA\EPLGSALELRASLEIDVAE\RGCE HGPSQQLPRCP*SWAWSEPWCQRPGCAV*APLP Y*REASFIYQSHSPAASGPFHSAGAGAVYLQAGG V/GEQEKEAVRKGSGSSSCSQRGP\PPPGMEVCPL **LGFWAICP** ASLSKPAGISTMPWALILLFLLTHSAVSVVQAGL 3411 1040 887 A TOPPSVSKDLR\QTATLTCTGNSNNVGHQGVIWL QQHQGHPPKLLSYRNNNRPSGISERLSAYKSGNA ASLTIYGLQTEHEAD**CRPRRKLIPKTARLFFFFL **IDNEEYLLRVY** 3412 RRGIPGSASLSLTMCVRSCFQSPRLQWVWRTAFL 164 83 A KHTORRHOGSHRWTHLGGSTYRAVIFDMGGVLI **PSPGRVAAEWEVQNRIPSGTILKALMEGGENGP** WMRFMRAEITAEGFLREFGRLCSEMLKTSVPVD SFFSLLTSERVAKQFPVMTEAITQIRAKGLQTAVL SNNFYLPNQKSFLPLDRKQFDVIVESCMEGICKP

000 10	Media	l Dungting	Drodieta 3	Amino seid seguence (AmAlonine Co-Cysteine Do-Associate Asia
SEQ ID NO:	Method	Predicted beginning nucleotide location corresponding to first amino acid residue of peptide	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion
-		sequence	·	DPRIYKLCLEQLGLQPSESIFLDDLGTNLKEAARL GIHTIKVNDPETAVKELEALLGFTLRVGVPNTRP VKKTMEIPKDSLQKYLKDLLGIQTTGPLELLQFD HGQSNPTYYIRLANRDLVLRKKPPGTLLPSAHAI EREFRIMKALANAGVPVPNVLDLCEDSSVIGTPF YVMEYCPGLIYKDPSLPGLEPSHRRAIYTAMNTV LCKIHSVDLQAVGLEDYGKQGSTTWV/YSSRRA RGALLFLDWELSYPWGDPFADVGYSCLAHYLPS SFPVLRGINDCDLTQLGIPAAEEYFRMYCLQMGL PPTENWNFYMAFSFFRVAAILQGVYKRSLTGQA SSTYAEQTGKLTEFVSNLAWDFAVKEGFRVFKE MPFTNPLTRSYHTWARPQSQWCPTGSRSYSSVPE ASPAHTSRGGLVISPESLSPPVRELYHRLKHFME QRVYPAEPELQSHQASAARWSPSPLIEDLKVKQP
3413	A	105	1573	W*GGRSGRTSWRLLALGCHT PESRHQCFSDRSSHFLTMEMEQEKMTMNKELSP DAAAYCCSACHGDETWSYNHPIRGRAKSRSLSA SPALGSTKEFRRTRSLHGPCPVTTFGPKACVLQN PQTIMHIQDPASQRLTWNKSPKSVLVIKKMRDAS LLQPFKELCTHLMEENMIVYVEKKVLEDPAIASD ESFGAVKKKFCTFREDYDDISNQIDFIICLGGDGT LLYASSLFQGSVPPVMAFHLGSLGFLTPFSFENFQ SQVTQVIEGNAAVVL/RGSRLKVRVVKELRGKK TAVHNGLGEKGSQAAGLDMDVGKQAMQYQVL NEVVIDRGPSSYLSNVDVYLDGHLITTVQGD/G* GPQHLSWGP*AFLGRE*RLRLSLSGVIVSTPTGST AYAAAAGASMIHPNVPAIMITPICPHSLSFRPIVV PAGVELKIMLSPEARNTAWVSFDGRKRQEIRHG DSISITTSCYPLPSICVRDPVSDWFESLAQCLHWN
3414	A	20	2602	VRKKQAHFEEEEEEEG VIVNKNVNWINYIYYNQQQRAFHELKEKLMSAL ALGLPDLTKPFTFYESEREKMAVGVLTQTVGPW PRPVAYLSKQLDGVSKGWPPCLRALAATALLAQ EADKLTLGQNLNIKAPHAVVTLMNTKGHHWLT NARLTKYQSLPCENPHITIEVCNTLNPTTLLPVSE SPGEHNCVEVLDSVYSSRPDLRDQPWASSVDWE LYMDGSSFINSQGERCAGYAVVTLDAVIKAKLW LQGTSAQKAELIALTRAVELSEGQESLEELLGRY FYVSHLPAFAKAVAQLCITCRQHNARQSPTVSPH IQAYGAAPFEDLQVDFTEMPKCGGNKYLLVLTC TYSGWVEAYPTRTEKAYEVTRVLLRDLIPRFGLP LRIGSHNGPVFVADLDCVEINVDTGVIWATWIKN EKDPVQLQKGKSGPSCTKGQCNPLELVITNPLDP RWKKGERVTLGINGAGLNPRVNILVRGEVYKCS LEPVFQTFYDELNVPITEFPGKTRNLFLQLAEHV AQSLTVTSCYVCGGTVIADQWPWEARELVPTDP VPDEFPAQKNHPDNFWVLKASIIRQYYIARVEKD FTLPVGRLHGG/RSNHTEKNPFSKFPKLQTV*AHP ESHRDWTAPTGLYWICGHRAYTKLP\ASSCVIGTI KPSFFLLSIKTGELLGFPVYASR\KSIAIRN*NNDK WPPERIIQYYGPAT*AQDGSWGYRIPIYMINRIIRL QAVLKIITATGRALTILAQQETQMRNAIYQNRLA LDYLLAAEGEVCRKFNLTNCCLHIDNQGQVVED IVRDMTKVAHVPVQVWHGFDPGAMFRKWFPAL GGFKTLIIRVIIVIGTYLLLPRLLPVLLQMIKSFIAT

SEQ ID	Method	Predicted	Predicted end	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid,
NO:		beginning	nucleotide	E-Glutamic Acid, F-Phenylalanine, G-Glycine, H-Histidine,
		nucleotide location	location corresponding	I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine,
		corresponding	to last amino	T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine,
		to first amino	acid residue of	X=Unknown, *=Stop codon, /=possible nucleotide deletion,
,		acid residue of peptide	peptide sequence	\=possible aucleotide insertion
		sequence	sequence	
				LVYQNASAQVYYINHY
3415	A	455	108	NMSWRGRSTYRPRPRRSLQPPELIGAMLEPTDEE
Ó				PKEEKPPTKSRNPTPDQKREDDSG/SAA*DFKWP EPGKPIFQGAMVRPKTGG/CGCEGGY*CQGEDS\P
				KAEHFKMPEAGEGKSQV
3416	A	1	874	FFFFQRINFIEHSGSVSLLALACDLGWCEDWSCC
				LVQGGGDLVDVVQTNHGEDEAGGDTDSVDEAR
	İ		i	CKESQQEAQENLREDLCLESFAKDKILQIIEGSER
				EHEETRTKQAALDGEPLGGGQLTAVHLHPSKEQ
ł	İ			QGQEGGERQRGARTHHWRGWEKGRRVRLRPPS
				GKLRADQPVRKLGGPTPS/TELPGLQPHAPTPHT
				A/PATPTYSPAPDTPNPPVRWKCPLPVEPRTRQLC
			1	RERTRKACPPKPRPPLGLPGDPTGPVTHHAPPVS PTGASGOERRAEPGAVSYAHASATK
3417	A	243	847	CLKYMYTYIFCPNCVSYKMKTDHFSLRYLHSSC
] 3717	A	273	347	AEDNKSSVDSSGQAAHPSKGKFFPHGTHWGTQC
				RGHISVLGWQCSCPSTGCRVGLGLAMCQTHAYI
				HTHTHTHTPTDYGAHHTDPLQRWGLGPR\KS
		ļ	l	EAGPLPQLSRDQSHPGPLSPGASPRSAGLPGWHP
:				AHQEPRARGRCARDGLSLQTRLTNKYDIQCCQE
				MRK
3418	Α	4073	1000	LDEYEARLTLANLDDFEEDNEDDDENRVNQEEK
				AAKITELINKLNFLDEAEKDLATVNSNPFDDPDA
				AELNPFGDPDSEEPITETASPRKTEDSFYNNSYNP FKEVQTPQYLNPFDEPEAFVTIKDSPPQSTKRKNI
				RPVDMSKYLYADSSKTEEEELDESNPFYEPKSTP
				PPNNLVNPVQELETERRVKRKAPAPPVLSPKTGV
:				LNENTVSAGKDLSTSPKPSPIPSPVLGRKPNASOS
,		J		LLVWCKEVTKNYRGVKITNFTTSWRNGLSFCAI
				LHHFRPDLIDYKSLNPQDIKENNKKAYDGFASIGI
				SRLLEPSDMVLLAIPDKLTVMTYLYQIRAHFSGQ
			ł	ELNVVQIEENSSKSTYKVGNYETDTNSSVDQEKF
				YAELSDLKREPELQQPISGAVDFLSQDDSVFVND
	1			SGVGESESEHQTPDDHLSPSTASPYCRRTKSDTEP
		J.	}	QKSQQSSGRTSGSDDPGICSNTDSTQAQVLLGKK RLLKAETLELSDLYVSDKKKDMSPPFICEETDEQ
				KLQTLDIGSNLEKEKLENSRSLECRSDPESPIKKT
				SLSPTSKLGYSYSRDLDLAKKKHASLRQTESDPD
				ADRTTLNHADHSSKIVQHRLLSRQEELKERARVL
				LEQARRDAALKAGNKHNTNTATPFCNRQLSDQ
	1		{	QDEERRRQLRERARQLIAEARSGVKMSELPSYGE
				MAAEKLKERSKASGDENDNIEIDTNEEIPEGFVV
		1		GGGDELTNLENDLDTPEQNSKLVDLKLKKLLEV
				QPQVANSPSSAAQKAVTESSEQDMKSGTEDLRT
		J		ERLOKTTERFRNPVVFSKDSTVRKTQLQSFSQYI
				ENRPEMKRQRSIQEDTKKGNEEKAAITETQRKPS EDEVLNKGFKDS\SQYVVGELAALENEQKQIDTR
				AALVEKRLRYLMDTGRNTEEEEAMMQEWFML
	1			VNKKNALIRRMNQLSLLEKEHDLERRYELLNRE
			1	LRAMLAIEDWQKTEAQKRREQLLLDELVALVN
			1	KRDALVRDLDAQEKQAEEEDEHLERTLEQNKG
	<u></u>		<u> </u>	KMAKKEEKCVLQ
3419	Α	4073	1000	LDEYEARLTLANLDDFEEDNEDDDENRVNQEEK
	}			AAKITELINKLNFLDEAEKDLATVNSNPFDDPDA
L		L	L	AELNPFGDPDSEEPITETASPRKTEDSFYNNSYNP

SEQ ID	Method	Predicted	Predicted end	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid,
NO:	Method	beginning	nucleotide	E-Glutamic Acid, F-Phenylalanine, G-Glycine, H-Histidine,
		nucleotide	location	I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine,
1		location corresponding	corresponding to last amino	T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine,
		to first amino	acid residue of	X=Unknown, *=Stop codon, /=possible nucleotide deletion,
		acid residue of	peptide	\=possible nucleotide insertion
		peptide sequence	sequence	
	 			FKEVQTPQYLNPFDEPEAFVTIKDSPPQSTKRKNI
				RPVDMSKYLYADSSKTEEEELDESNPFYEPKSTP
	1	1		PPNNLVNPVQELETERRVKRKAPAPPVLSPKTGV
		}		LNENTVSAGKDLSTSPKPSPIPSPVLGRKPNASQS
				LLVWCKEVTKNYRGVKITNFTTSWRNGLSFCAI
				LHHFRPDLIDYKSLNPQDIKENNKKAYDGFASIGI SRLLEPSDMVLLAIPDKLTVMTYLYQIRAHFSGQ
				ELNVVQIEENSSKSTYKVGNYETDTNSSVDQEKF
				YAELSDLKREPELQQPISGAVDFLSQDDSVFVND
				SGVGESESEHQTPDDHLSPSTASPYCRRTKSDTEP
				QKSQQSSGRTSGSDDPGICSNTDSTQAQVLLGKK
				RLLKAETLELSDLYVSDKKKDMSPPFICEETDEQ
				KLQTLDIGSNLEKEKLENSRSLECRSDPESPIKKT
				SLSPTSKLGYSYSRDLDLAKKKHASLRQTESDPD
ľ	ļ		ĺ	ADRTTLNHADHSSKIVQHRLLSRQEELKERARVL
				LEQARRDAALKAGNKHNTNTATPFCNRQLSDQ
				QDEERRRQLRERARQLIAEARSGVKMSELPSYGE
				MAAEKLKERSKASGDENDNIEIDTNEEIPEGFVV GGGDELTNLENDLDTPEQNSKLVDLKLKKLLEV
	}			QPQVANSPSSAAQKAVTESSEQDMKSGTEDLRT
				ERLOKTTERFRNPVVFSKDSTVRKTQLQSFSQYI
ĺ				ENRPEMKRQRSIQEDTKKGNEEKAAITETQRKPS
			İ	EDEVLNKGFKDS\SQYVVGELAALENEQKQIDTR
ļ				AALVEKRLRYLMDTGRNTEEEEAMMQEWFML
		'	İ	VNKKNALIRRMNQLSLLEKEHDLERRYELLNRE
				LRAMLAIEDWQKTEAQKRREQLLLDELVALVN
				KRDALVRDLDAQEKQAEEEDEHLERTLEQNKG
<u> </u>	ļ			KMAKKEEKCVLQ
3420	A	612	1058	ENLGPNYSHRLLHHPTFYKKIHKKHHEWTAPIG VISLYAHPIEHAVSNMLPVIVGPLVMGSHLSSITM
ł				WFSLALIITTISHCGYHLPFLPSPEFHDYHHLKFN
				QCYGVLGVLDHLHGTDTMFKQTKAYERHVLLL
		1		GFTPLSESIPDSPK
3421	A	23	2005	LLTPCDGRIPGRPSVGAESGSDFQQRRRRRRDPE
3 121			2000	EPEKTELSERELAVAVAVSQENDEENEERWVGP
				LPVEATLAKKRKVLEFERVYLDNLPSASMYERS
				YMHRDVITHVVCTKTDFIITASHDGHVKFWKKIE
				EGIEFVKHFRSHLGVIESIAVSSEGALFCSVGDDK
				AMKVFDVVNFDMINMLKLGYFPGQCEWIYCPG
				DAISSVAASEKSTGKIFIYDGRGDNQPLHIFDKLH
				TSPLTQIRLNPVYKAVVSSDKSGMIEYWTGPPHE
				YKFPKNVNWEYKTDTDLYEFAKCKAYPTSVCFS PDGKKIATIGSDRKVRIFRFVTGKLMRVFDESLS
		!		MFTELQQMRQQLPDMEFGRRMAVERELEKVDA
		1	1	VRLINIVFDETGHFVLYGTMLGIKVINVETNRCV
	1	1		RILGKQENIRVMQLALFQGIAKKHRAATTIEMKA
				SENPVLQNIQADPTIVCTSFKKNRFYMFTKREPE
				DTKSADSDRDVFNEKPSKEEVMAATQAEGPKRV
				SDSAIIHTSMGDIHTKLFPVECPKTVENFCVHSRN
				GYYNGHTFHRIIKGFMIQTGDPTGTGMGGESIWG
				GEFEDEFHSTLRHDRPYTLSMANAGSNTNGSQFF
	1	1		ITVVPTPWLDNKHTVFGRVTKGMEVVQRISN\VK
				VNPKTDKPYEDVSIINITVK
3422	Α	2486	433	FVLVCAPLTWAGARHRMAASKKPPRVRVNHQ
	L		L	DFQLRNLRIIEPNEVTHSGDTGVETDGRMPPKVT

Method	Predicted beginning nucleotide location corresponding to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion
			SELLRQLRQAMRNSEYVTEPIQAYIIPSGDAHQSE YIAPCDCRRAFVSGFDGSAGTAIITEEHAAMWTD GRYFLQAAKQMDSNWTLMKMGLKDTPTQEDW LVSVLPEGSRVGVDPLIIPTDYWKKMAKVLRSA GHHLIPVKENLVDKIWTDRPERPCKPLLTLGLDY TGISWKDKVADLRLKMAERNVMWFVVTALDEI AWLFNLRGSDVEHNPVFFSYAIIGLETIMLFIDGD RIDAPSVKEHLLLDLGLEAEYRIQVHPYKSILSEL KALCADLSPREKVWVSDKASYAVSETIPKDHRC CMPYTPICIAKA\VKNSA\ESEGMRRAHIKDAVAL CELFNWLEKEVPKGGVTEISAADKAEEFRRQQA DFVDLSFPTISSTGPNGAIIHYAPVPETNRTLSLDE VYLIDSGAQYKDGTTDVTRTMHFGTPTAYEKEC FTYVLKGHIAVSAAVFPTGTKGHLLDSFARSAL WDSGLDYLHGTGHGVGSFLNVHEGPCGISYKTF SDEPLEAGMIVTDEPGYYEDGAFGIRIENVVLVV PVKTKYNFNNRGSLTFEPLTLVPIQTKMIDVDSL TDKECDWLNNYHLTCRDVIGKELQKQGRQEAL
A	5515	934	EWLIRETQPISKQH FKMPENPATDKLQVLQVLDRLKMKLQEKGDTS QNEKLSMFYETLKSPLFNQILTLQQSIKQLKGQL NHIPSDCSANFDFSRKGLLVFTDGSITNGNVHRPS NNSTVSGLFPWTPKLGNEDFNSVIQQMAQGRQIE YIDIERPSTGGLGFSVVALRSQNLGKVDIFVKDV QPGSVADRDQRLKENDQILAINHTPLDQNISHQQ AIALLQQTTGSLRLIVAREPVHTKSSTSSSLNDTT LPETVCWGHVEEVELINDGSGLGFGIVGGKTSGV VVRTIVPGGLADRDGRLQTGDHILKIGGTNVQG MTSEQVAQVLRNCGNSVRMLVARDPAGDISVTP PAPAALPVALPTVASKGPGSDSSLFETYNVELVR KDGQSLGIRIVGYVGTSHTGEASGIYVKSIIPGSA AYHNGHIQVNDKIVAVDGVNIQGFANHDVVEVL RNAGQVVHLTLVRRKTSSSTSPLEPPSDRGTVVE PLKPPALFLTGAVETETNVDGEDEEIKERIDTLKN DNIQALEKLEKVPDSPENELKSRWENLLGPDYEV MVATLDTQIADDAELQKYSKLLPHTLRLGVEV DSFDGHHYISSIVSGGPVDTLGLLQPEDELLEVN GMQLYGKSRREAVSFLKEVPPPFTLVCCRRLFDD EASVDEPRRTETSLPETEVDHNMDVNTEEDDDG ELALWSPEVKIVELVKDCKGLGFSILDYQDPLDP TRSVIVIRSLVADGVAERSGGLLPGDRLVSVNEY CLDNTSLAEAVEILKAVPPGLVHLGICKPLVEDN EEESCYILHSSSNEDKTEFSGTIHDINSSLILEAPK GFRDEPYFKEELVDEPFLDLGKSFHSQQKEIEQS KEAWEMHEFLTPRLQEMDEEREMLVDEEYELY QDPSPSMELYPLSHIQEATPVPSVNELHFGTQWL HDNEPSESQEARTGRTVYSQEAQPYGYCPENVM KENFVMESLPSVPSTEGNSQQGRFDDLENLNSLA KTSLDLGMIPNDVQGPSLLIDLPVVAQRREQEDL PLYQHQATRVISKASAYTGMLSSRYATDTCELPE REEGEGEETPNFSHWGPPRIVEIFREPNVSLGISIV GGQTVIKRLKNGEELKGIFIKQVLEDSPAGKTNA LKTGDKILEVSGVDLQNASHSEAVEAIKNAGNP VVFIVQSLSSTPRVIPNVHNKANKITGNQNQDTQ
		beginning nucleotide location corresponding to first amino acid residue of peptide sequence	beginning nucleotide location corresponding to first amino acid residue of peptide sequence

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SEQ ID	Method	Predicted	Predicted end	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid,
NO:		beginning	nucleotide	E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methlonine,
		nucleotide location	location corresponding	I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine,
		corresponding	to last amino	T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine,
	ļ	to first amino	acid residue of	X=Unknown, *=Stop codon, /=possible nucleotide deletion,
		acid residue of	peptide	∖=possible nucleotide insertion
		peptide sequence	sequence	
				DAFTDQKIRQRYADLPGELHIIELEKDKNGLGLS LAGNKDRSRMSIFVVGINPEGPAAADGRMHIGD
1				ELLEINNQILYGRSHQN\ASAIIKTAPSKVKLVFIR
ł		-0		NEDAVNQMAVTPFPVPSSSPSSIEDQSGTEPISSEE
! .•				\DGSLE\VGIKQLPESESFKLAVSQMKQQKYPTKV
			1	SFSSQEIPLAPASSYHSTDADFTGYGGFQAPLSVD
		i]	PATCPIVPGQEMIIEISKRRSGLGLSIVGGKDTPLV
				NGVDLRNSSHEEAITALRQTPQKVRLVVYRDEA
				HYRDEENLEIFPVDLQKKAGRGLGLSIVGKR
3424	A	2223	1162	HASERVVQLPDFVWDQYTHSLGRVEREFKNRKR
				HTRRVKLVFDKGLPARPKSPLDPKKDGESLSYS
				MLPLSDGPEGSSSRPQMIRGRLCDDTKPETFNQL
				WTVEEQKKLEQLLIKYPPEEVESRRWQKIADELG
				NRTAKQVASRVQKYFIKLTKAGIPVPGRTPNLYI
		ļ.		YSKKSSTSRRQHPLNKHLFKP\GTFMTSHEPPVY
				MDEDDDRSCFHSHMNTAVEDASDDESIPIMYRN
	E			LPEYKELLQFKKLKKQKLQHMQAESGFVQHVGF
	Į			KCDNCGIEPIQG\VRW\HCR\DCPP\EMSL\DFC\DS
				C\SDCLHET\DIHKGDHQLEPIYRS\ETFLDRDYCV
				SQGTSYNYLDPNYFPANR
3425	A	2223	1162	HASERVVQLPDFVWDQYTHSLGRVEREFKNRKR
				HTRRVKLVFDKGLPARPKSPLDPKKDGESLSYS
				MLPLSDGPEGSSSRPQMIRGRLCDDTKPETFNQL
ļ				WTVEEQKKLEQLLIKYPPEEVESRRWQKIADELG
				NRTAKQVASRVQKYFIKLTKAGIPVPGRTPNLYI
				YSKKSSTSRRQHPLNKHLFKP\GTFMTSHEPPVY
				MDEDDDRSCFHSHMNTAVEDASDDESIPIMYRN
				LPEYKELLQFKKLKKQKLQHMQAESGFVQHVGF
•				KCDNCGIEPIQG\VRW\HCR\DCPP\EMSL\DFC\DS
				C\SDCLHET\DIHKGDHQLEPIYRS\ETFLDRDYCV
			1.22	SQGTSYNYLDPNYFPANR
3426	A	2	1553	LFVVVHDDPRWGTPRYWLGALYRNQQSSPTAPP
		1,		GLLPLEYFPAAPHCSHSRQWRCSQTHRIHHHPQ
				MLGPCRQEICGITMAAGTLYTYPENWRAFKALI
•	· I			AAQYSGAQVRVLSAPPHFHFGQTNRTPEFLRKFP
				AGKVPAFEGDDGFCVFESNAIAYYVSNEELRGST
				PEAAAQVVQWVSFADSDIVPPASTWVFPTLGIM
				HHNKQATENAKEEVRRILGLLDAYLKTRTFLVG
				ERVTLADITVVCTLLWLYKQVLEPSFRQAFPNTN
				RWFLTCINQPQFRA\VFGEVKLCEKMAQF\DAKK
			1	FAETQPKKDTPRKEKGSREEKQKPQAERKEEKK
	1			AAAPAPEEEMDECEQALAAEPKAKDPFAHLPKS
		1	1	TFVLDEFKRKYSNEDTLSVALPYFWEHFDKDGW
				SLWYSEYRFPEELTQTFMSCNLITGMFQRLDKLR
		1		KNAFASVILFGTNNSSSISGVWVFRGQELAFPLSP
		1		DWQVDYESYTWRKLDPGSEETQTLVREYFSWE
				GAFQHVGKAFNQGKIFK
3427	A	755	52	TAARRQKGTAARRQKGTAARR
	-	1		RQKGTAARRRQKGTAARRRQKGT
1	1	1		AARRQKGTAARRQKGTAARRR
				QKGLSNLDAAEWLPPKKG\GEKKKGPFLAINEV
				VT\REYPINILKRIHGVGFKKRAPRALKEIRKFAM
				KEMGTPDVRIDTRLNKAVWAKGIRNVPYRIRVR
		1		LSRKRNEDEDSPNKLYTLVTYVPVTTFKNLQTV
ļ	1	L	J	NVDEN

SEQ ID NO:	Method	Predicted beginning nucleotide location corresponding to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion
3428	A	4	1939	LPLSLSFSEMPLPLLPMDLKGEPGPPGKPGPWGP PGPPGFPGKPGHGKPGLHGQPGPAGPPGFSRMG KAGPPGLPGNVGPPGQPGLRGEPGIRGDQGLRGP PGPPGLPGPSGITIPGKPGAQGVPGPPGFQGEPGP QGEPGPPGDRGLKGDNGVGQPGLPGAPGQGGAP GPPGLPGPAGLGKPGLDGLPGAPGDKGESGPPG VPGPRGEPGAVGPKGPPGVDGVGVPGAAGLPGP QGPSGAKGEPGTRGPPGLIGPTGYGMPGLPGPKG DRGPAGVPGLLGDRGEPGEDGEPGEQGPQGLGG PPGLPGSAGLPGRRGPPGPKGEAGPGGPPGVPGI RGDQGPSGLAGKPGVPGERGLPGAHGPPGPTGP KGEPGFTGRPGGPGVAGALGQKGDLGLPGQPGL RGPSGIPGLQGPAGPIGPQGLPGLKGEPGLPGPPG EGRAGEPGTAGP\RGPPGVPGSPGITGPPG\LPGPP GAPGAFDETGIAGLHLPNGGVEGAVLGKGGKPQ FGLGELSAHATPAFTAVLTSPLPASGMPVKFDRT LYNGHSGYNPATGIFTCPVGGVYYFAYHVHVKG TNVWVALYKNNVPATYTYDEYKKGYLDQASG GAVLQLRPNDQVWVQMPSDQANGLYSTEYIHSS FSGFLLCPT
3429	A	212	1075	EGLTGPCERVPFLLGRGPPHGATRAGHRRAVRW AGPESLPPLPRSLIMDSPRAGTHQGPLDAETEVG ADRCTSTAYQEQRPQVEQVGKQAPLSPGLPAMG GPGPGPCEDPAGAGGAGAGGSEPLVTVTVQCAF TVALRARRGADLSSLRALLGQALPHQ\AQLGQLS YLAPGEDGHWVPIPEEESLQRAWQDAAACPRGL QLQCRGAGGRPVLYQVVAQHSYSAQGPEDLGF RQGDTVDVLCEVDQAWLEGHCDGRIGIFPKCFV VPAGPRMSGAPGRLPRSQQGDQP
3430	A .	799	1989	INKYINIRKKIKLLSPLPPLWSHLALLQASATKWV LTPAAFAGKLLSVFRQPLSSLWRSLVPLFCWLRA TFWLLATKRRKQQLVLRGPDETKEEEEDPPLPTT PTSVNYHFTRQCNYKCGFCFHTAKTSFVLPLEEA KRGLLLLK\EAG\LEKINFSGG\EPFLQDRGEYLGK LVRFCKVELRLPSVSI\VSNGSLIRERWFQNYG\E YLDILAISCDSFDEEVNCP\IGRGN\GKKNHVENL QKL\RRWCRDYRVPFKINSVINPF\NVEEDMTEQI KALNPVRWKVFQCLLIEGENCGEDA\LREAERFV IGDEEFERFLERHKEVSCLVPESNQKMKDSYLIL DEYMRFLNCRKGRKDPSKSILDVGVEEAIKFSGF DEKMFLKRGGKYIWSKADLKLDW
3431	A	5468	2146	ACGFLPGRCHFSTFKQCQEWLSRLSRATARPAKP EDLFAFAYHAWCLGLTEEDQHTHLCQPGEHIRC RQEAELARMGFDLQNVWRVSHINSNYKLCPSYP QKLLVPVWITDKELENVASFRSWKRIPVVVYRH LRNGAAIARCSQPEISWWGWRNADDEYLVTSIA KACALDPGTRATGGSLSTGNNDTSEACDADFDS SLTACSGVESTAAPQKLLILDARSYTAAVANRAK GGGCECEEYYPNCEVVFMGMANIHAIRNSFQYL RAVCSQMPDPSNWLSALESTKWLQHLSVMLKA AVLVANTVDREGRPVLVHCSDGWDRTPQIVALA KILLDPYYRTLEGFQVLVESDWLDFGHKFGDRC GHQENVEDQNEQCPVFLQWLDSVHQLLKQFPCL FEFNEAFLVKLVQHTYSCLYGTFLANNPC\EREK RNIYK/RGTCSVWALLRAGNKNFHNFLYTPSSD

SEQ ID Method Predicted Predicted end Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, NO: beginning nucleotide I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, nucleotide location N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, corresponding location corresponding T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, to last amino X=Unknown, *=Stop codon, /=possible nucleotide deletion, acid residue of to first amino \=possible nucleotide insertion acid residue of peptide peptide sequence sequence MVLHPVCHVRALHLWTAVYLPASSPCTLGEEN **MDLYLSPVAQSQEFSGRSLDRLPKTRSMDDLLS** ACDTSSPLTRTSSDPNLNNHCQEVRVGLEPWHS NPEGSETSFVDSGVGGPQQTVGEVGLPPPLPSSQ KDYLSNKPFKSHKSCSPSYKLLNTAVPREMKSNT SDPEIKVLEETKGPAPDPSAQDELGRTLDGIGEPP **EHCPETEAVSALSKVISNKCDGVCNFPESSQNSPT** GTPQQAQPDSMLGVPSKCVLDHSLSTVCNPPSA ACQTPLDPSTDF\LNQDPSGSVASISHQEQLSSVP DLTHGEEDIGKRGNNRNGOLLENPRFGKMPLEL VRKPISQSQISEFSFLGSNWDSFQGMVTSFPSGEA TPRRLLSYGCCSKRPNSKQMRATGPCFGGQWAQ REGVKSPVCSSHSNGHCTGPGGKNQMWLSSHPK QVSSTKPVPLNCPSPVPPLYLDDDGLPFPTDVIQH RLRQIEAGYKQEVEQLRRQVRELQMRLDIRHCC APPAEPPMDYEDDFTCLKESDGSDTEDFGSDHSE DCLSEASWEPVDKKETEVTRWVPDHMASHCYN CDCEFWLAKRRHHCRNCGNVFCAGCCHLKLPIP DQQLYDPVLVCNSCYEHIQVSRARELMSQQLKK **PIATASS** MTFFSSVADFIGLDPRIAAWLIDPSDATPSFEDLV 3432 36 1873 A EKYCEKSITVKVNSTYGNSSRNIVNQNVRENLKT LYRLTMDLCSKLKDYGLWQLFRTLELPLIPILAV MESHAIQVNKEEMEKTSALLGARLKELEQEAHF VAGERFLITSNNQLREILFGKLKLHLLSQRNSLPR TGLQKYPSTVSEALNALRDLHPLPKIILEYRQVH KIKSTFVDGLLACMKKGSISSTWNQTGTVTGRLS AKHPNIQGISKHPIQITTPKNFKGKEDKILTISPRA MFVSSKGHTFLAADFSQIELRILTHLSGDPELLKL **FQESERDDVFSTLTSQWKDVPVEQVTHADREQT** KKVVYAVVYGAGKERLAACLGVPIQEAAQFLES FLOKYKKIKDFARAAIAOCHOTGCVVSIMGRRR PLPRIHAHDQQLRAQAERQAVNFVVQGSAADLC KLAMIHVFTAVAASHTLTARLVAQIHDELLFEVE DPQIPECAALVRRTMESLEQVPLKVSLSAGRSWG HLVPLQEAW\ALRQAHVALSLPATAWLPLGPLP **APSPHPCIFRLHFVCSPRQQWEERTGFQQSIVWPS** PRSPALYAPGRINPLGLGWPAIPWSKCLCKALKK **IPPKERAPGIRASCLAITAGARPTSYGRVGCEGDV** 3433 Ā 1481 476 RLSPVSPLLAPPDPRLASRWEGRSRMKGKKGIVA ASGSETEDEDSMDIPLDLSSSAGSGKRRRRGNLP KESVQILRDWLYEHRYNAYPSEQEKALLSQQTH LSTLQVCNWFINARRRLLPDMLRKDGKDPNQFTI SRRGAKISETSSVESVMGIKNFMPALEETPFHSFT\ AGPNPTLG\RPLSAKP/SQSPGSVLARPSVICHTTV TAIERLSLSCQSVGCGQNT\DIQQIAT\RNLRDS SLMYPEDTCKSGPSTNTQSGLFNTPPPTPPDLNQ DFSGFQLLVDVALKRAAEMELQAKLTA 3434 A 1720 1243 NGPVPPGGSKTKWAGGSAAEGSPRLSPSPGAAQ VPALLRGEPRGGAAAGSFWKPLHQHSCGLRPPP/ PPD/RLSRLPGKTLSACDRENGARRPLLLGSTSFIP **IGRRTYASAAEPVGSKAVLVTGCDSGFGFSLAKH** LHSKGFLVFAGCLMKDKGHDGVKELDSLNSDRL RTVQLNVCSSEEVEKV/VGDCPLEPEGP\EKGMW

SEO ID Method Predicted Predicted end Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, nucleotide NO: beginning E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, nucleotide location I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, corresponding N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, location to last amino T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, corresponding X=Unknown, *=Stop codon, /=possible nucleotide deletion, acid residue of to first amino acid residue of peptide \=possible nucleotide insertion peptide sequence sequence GLVNNAGISTFGEVEFTSLETYKQVAEVNLWGT VRMTKSFLPLIRRAKGRVVNISSMLGRMANPAR SPYCITKFGVEAFSDCLRYEMYPLGVKVSVVEPG NFIAATSLYSPESIQAIAKKMWEELPEVVRKDYG KKYFDEKIAKMETYCSSGSTDTSPVIDAVTHALT ATTPYTRYHPMDYYWWLRMQIMTHLPGAISDM ENQQOMLVAKEQRLHFLKQQERRQQQSISENEK 3435 Α 842 3595 LOKLKERVEAQENKLKKIRAMRGQVDYSKIMN GNLSAEIERFSAMFQEKKQEVQTAILRVDQLSQQ LEDLKKGKLNGFOSYNGKLTGPAAVELKRLYOE LQIRNQLNQEQNSKLQQQKELLNKRNMEVAMM DKRISELRERLYGKKIQACEKVFLNRVNGTSSPQ **SPLSTSGRVAAVGPYIQVPSAGSFPVLGDPIKPQS** LSIASNAAHGRSKSANDGNWPTLKQNSSSSVKP VQVAGADWKDPSVEGSVKQGTVSSQPVPFSALG **PTEKPGIEIGKVPPPIPGVGKQLPPSYGTYPSPTPL** GPGSTSSLERRKEGSLPRPSAGLPSRQRPTLLPAT GSTPQPGSSQQIQQRISVPPSPTYPPAGPPAFPAGD SKPELPLTVAIRPFLADKGSRPQSPRKGPQTVNSS SIYSMYLQQATPPKNYQPAAHSALNKSVKAVYG KPVLPSGSTSPSPLPFLHGSLSTGTPQPQPPSESTE KEPEQDGPAAPADGSTVESLPRPLSPTKLTPIVHS PLRYQSDADLEALRRKLANAPRPLKKRSSITEPE GPGGPNIQKLLYQRFNTLAGGMEGTPFYQPSPSQ DFMVTLADVDNGNTNANGNLEELPPAQPTAPLP AEPAPSSDANDNELPSPEPEELICPQTTHQTAEPA **EDNNNNVATVPTTEQIPSPVAEAPSPGEEQVPPA** PLPPASHPPATSTNKRTNLKKPNSERTGHGLRVR FNPLALLLDASLEGEFDLVQRIIYEVEDPSKPNDE GITPLHNAVCAGHHHIVKFLLDFGVNVNAADSD **GWTPLHCAASCNSVHLCKQLVESGAAIFASTISD IETAADKCEEMEEGYIQCSQFLYGVQEKLGVMN** KGVAYALWDYEAQNSDELSFHEGDALTILRRKD 3436 3 2604 **GSTHASEKMKTGRSALVVTDTGDMSVLNSPRHQ** Α SCIMHVDMDCFFVSVGIRNRPDLKGKPVAVTSN RGTGRAPLRPGANPQLEWQYYQNKILKGKADIP DSSLWENPDSAQANGIDSVLSRAEIASCSYEARQ LGIKNGMFFGHAKOLCPNLQAVPYDFHAYKEVA QTLYETLAS\YTHNIEAVSCDEALVDITEILAETK LTPDEFANAVRMEIKDQTKCAASVGIGSNILLAR MATRKAKPDGQYHLKPEEVDDFIRGQLVTNLPG VGHSMESKLASLGIKTCGDLQYMTMAKLQKEF **GPKTGQMLYRFCRGLDDRPVRTEKERKSVSAEI** NYGIRFTQPKEAEAFLLSLSEEIQRRLEATGMKG KRLTLKIMVRKPGAPVETAKFGGHGICDNIARTV TLDQATDNAKIIGKAMLNMFHTMKLNISDMRGV **GIHVNQLVPTNLNPSTCPSRPSVQSSHFPSGSYSV** RDVFQVQKAKKSTEEEHKEVFRAAVDLEISSASR TCTFLPPFPAHLPTSPDTNKAESSGKWNGLHTPV SVQSRLNLSIEVPSPSQLDQSVLEALPPDLREQVE QVCAVQQAESHGDKKKEPVNGCNTGILPQPVGT VLLQIPEPQESNSDAGINLIALPAFSQVDPEVFAA

LPAELQRELKAAYDQRQRQGENSTHQQSASASV

SEQ ID NO:	Method	Predicted beginning nucleotide location corresponding to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methlonine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \text{\text{=-possible}} nucleotide insertion
				PKNPLLHLKAAVKEKKRNKKKKTIGSPKRIQSPL NNKLLNSPAKTLPGACGSPQKLIDGFLKHEGPPA EKPLEELSASTSGVPGLSSLQSDPAGCVRPPAPNL AGAVEFNDVKTLLREWITTISDPMEEDILQVVKY CTDLIEEKDLEKLDLVIKYMKRLMQQSVESVWN MAFDFILDNVQVVLQQTYGSTLKVT
3437		32	4038	SLLRLLKAQWGSSGAASEPVVLGEEGCGFPSTNE YPDLEEERATYPQEEDRFLTPGRAQLLWSPWSPL DQEEACASRQLHSLASFSTVTARRNPLHNPWGM ELAASENTDSPSPRPLRPGVTLPPGALTMNTKDT TEVAENSHHLKIFLPKKLLECLPRCPLLPPERLRW NTNEEIASYLITFEKHDEWLSCAPKTRPQNGSIIL YNRKKVKYRKDGYLWKKRKDGKTTREDHMKL KVQGMECLYGCYVHSSIVPTFHRRCYWLLQNPD IVLVHYLNVPALEDCGKGCSPIFCSISSDRREWLK WSREELLGQLKPMFHGIKWSCGNGTEEFSVEHL VQQILDTHPTKPAPRTHACLCSGGLGSGSLTHKC SSTKHRIISPKVEPRALITLTSIPHPHPPEPPPLIAPLP PELPKAHTSPSSSSSSSSSGFAEPLEIRPSPPTSRGG SSRGGTAILLLTGLEQRAGGLTPTRHLAPQADPR PSMSLAVVVGTEPSAPPAPPSPAFDPDRFLNSPQR GQTYGGGQGVSPDFPEAEAAHTPCSALEPAAAL EPQAAARGPPPQSVAGGRRGNCFFIQDDDSGEEL KGHGAAPPIPSPPPSPPSPAPLEPSSRVGRGEALF GGPVGASELEPFSLSSFPDLMGELISDEAPSIPAPT PQLSPALSTITDFSPEWSYPEGGVKVLITGPWTEA AEHYSCVFDHIAVPASLVQPGVLRCYCPAHEVG LVSLQVAGREGPLSASVLFEYRARRFLSLPSTQL DWLSLDDNQFRMSILERLEQMEKRMAEIAAAGQ VPCQGPDAPPVQDEGQGPGFEARVVVLVESMIP RSTWKGPERLAHGSPFRGMSLLHLAAAQGYARL IETLSQWRSVETGSLDLEQEVDPLNVDHFSCTPL MWACALGHLEAAVLLFRWNRQALSIPDSLGRLP LSVAHSRGHVRLARCLEELQRQEPSVEPPFALSP PSSSPDTGLSSVSSPSELSDGTFSVTSAYSSAPDGS PPPAPLPASEMTMEDMAPGQLSSGVPEAPLLLM DYEATNSKGPLSSLPALPPASDDGAAPEDADSPQ AVDVIPVDMISLAKQIIEATPERIKREDFVGLPEA GASMRERTGAVGLSETMSWLASYLLENVDHFPS STPPSELYPFER\GRLGLSLTAPSWAEFLSCIPPVGK IGKLIFALLTL\SD\QEQRELYEAARVIQTAFRKYK GRRLKEQQEVAAAVIQRCYRKYKQLTWIALKFA LYKKMTQAAILIQSKFRSYYEQKRFQQSRRAAV LIQQHYRSYRRRPGPPHRTSATLPARNKGSFLTK KQDQAARKIMRFLRRCRHRMRELKQNQELEGLP OPGLAT
3438	A	469	2602	FGRLLWGTAFKSWKMKAPIPHLILLYATFTQSLK VVTKRGSADGCTDWSIDIKKYQVLVGEPVRIKC ALFYGYIRTNYSLAQSAGLSLMWYKSSGPGDFE EPIAFDGSRMSKEEDSIWFRPTLLQDSGLYACVIR NSTYCMKVSISLTVGENDTGLCYNSKMKYFEKA ELSKSKEISCRDIEDFLLPTREPEILWYKECRTKT WRPSIVFKRDTLLIREVREDDIGNYTCELKYGGF VVRRTTELTVTAPLTDKPPKLLYPMESKLTIQET QLGDSANLTCRAFFGYSGDVSPLIYWMKGEKFIE

SEQ ID NO:	Method	Predicted beginning nucleotide location corresponding to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion
		sequence		DLDENRVWESDI\KILKEHLGEQEVSISLIVDSVEE GDLGNYSCYVENGNGRRHASVLLHKRELMYTV ELAGGLGAILLLLVCLVTTYKCYKIEIMLFYRNHF GAEELDGDNKDYDAYLSYTKVDPDQWNQETGE
				EERFALEILPDMLEKHYGYKLFIPDRDLIPTGTYI EDVARCVDQSKRLIIVMTPNYVVRRGWSIFELET RLRNMLVTGEIKVILIECSELRGIMNYQEVEALK HTIKLLTVIKWHGPKCNKLNSKFWKRLQYEMPF
				KRIEPITHEQALDVSEQGPFGELQTVSAISMAAAT STALATAHPDLRSTFHNTYHSQMRQKHYYRSYE YDVPPTGTLPLTSIGNQHTYCNIPMTLINGQRPQT KSSREQNPDEAHTNSAILPLLPRETSISSVIW
3439	A	251	2037	GPGNSSILIGGGHLFLIRSCLNLLLLNSKENTEHT MAKKVAVIGAGVSGLSSIKCCVDEDLEPTCFERS DDIGGLWKFTERGSSLSVMIWPLALSLLRHGGFC
				YSDFPFHEDYPNFMNHEKFWDYLQEFAEHFDLL KYIQFKTTVCGITKRPDFSETGQWDVVTETEGKQ NRAVFDAVMVCTGHFLNPHLPLEAFPGIHKFKG QILHSQEYKIPEGFQGKRVLVIGLGNTGGDIAVEL
				SRTAAQVLLSTRTGTWVLGRSSDWGYPYNMMV TRRCCSFIAQVLPSRFLNWIQERKLNKRFNHEDY GLSITKGKKAKFIVNDELPNCILCGAITMKTSVIE FTETSAVFEDGTVEENIDVVIFTTGYTFSFPFFEEP
				LKSLCTKKIFLYKQVFPLNLERATLAIIGLIGLKGS ILSGTELQARWVTRVFKGLCKRPASQKLMMEAT EKEQLIKRGVFKDTSKDKFDYIAYMDDIAACIGT
				KPSIPLLFLKDPRLAWEVFFGPCTPYQYR\LMGPG KWDGARNAILTQWDRTLKPLKTRIVPDSSKAWP SM\SHYLKAWGAPVLLASLLLICK\SSLFLKLVRD KLQDRMSPYLVSLWRG
3440	A	1	3533	IMPCGSSRLLRGCWTHPNEPVSDLSYFDCIESVM ENSKVLGESMAGISQNAKTGDLPAFGECVGIASK ALCGLTEAAAQAAYLVGIFDPNSQAGHQGLVDP
				IQFARANQAIQMACQNLVDPGSSPSQVLSAATIV AKHTSALCNACRIASSKTANPVAKRHFVQSAKE VANSTANLVKTIKALDGDFSEDNRNKCRIATAPL
:				IEAVENLTAFASNPEFVSIPAQISSEGSQAQEPILV SAKPMLESSSYLIRTARSLAINPKDPPTWSVLAG HSHTVSDSIKSLITSIRDKAPGQRECDYSIDGINRC IRDIEQASLAAVSQSLATRDDISVEALQEQLTSVV
				QEIGHLIDPIATAARGEAAQLGHKGTQLASYFEP LILAAVGVASKILDHQQQMTVLDQTKTLAESAL QMLYAAKEGGGNPKAQHTHDAITEAAQLMKEA
				VDDIMVTLNEAASEVGLVGGMVDAIAEAMSKL DEGTPPEPKGTFVDYQTTVVKYSKAIAVTAQEM MTKSVTNPEELGGLASQMTSDYGHLAFQGQMA
				AATAEPEEIGFQIRTRVQDLGHGCIFLVQKAG\AL QVCPTDSYTKRELIECARAVTEKVSLVLSALQAG NKGTQACITAATAVSGIIADLDTTIMFATAGTLN AENSETFADHRENILKTAKALVEDTKLLVSGAAS
				TPDKLAQAAQSSAATITQLAEVVKLGAASLGSD DPETQVVLINAIKDVAKALSDLISATKGAASKPV DDPSMYQLKGAAKVMVTNVTSLLKTVKAVEDE
				ATRGTRALEATIECIKQELTVFQSKDVPEKTSSPE ESIRMTKGITMATAKAVAAGNSCRQEDVIATAN

.........

SEO ID Method Predicted end Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, Predicted NO: beginning nucleotide E-Glutamic Acid, F-Phenylalanine, G-Glycine, H-Histidine, location I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, nucleotide corresponding N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, location corresponding to last amino T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, to first amino acid residue of \=possible nucleotide insertion acid residue of peptide peptide sequence sequence LSRKAVSDMLTACKQASFHPDVSDEVRTRALRF GTECTLGYLDLLEHVLVILQKPTPELKQQLAAFS KRVAGAVTELIQAAEAMKGTEWVDPEDPTVIAE TELLGAAASIEAAAKKLEQLKPRAKPKQADETL DFEEQILEAAKSIAAATSALVKSASAAQRELVAQ **GKVGSIPANAADDGQWSQGLISAARMVAAATSS** LCEAANASVQGHASEEKLISSAKQVAASTAQLL VACKVKADQDSEAMRRLQAAGNAVKRASDNL VRAAQKAAFGKADDDDVVVKTKFVGGIAQIIAA QEEMLKKERELEEARKKLAQIRQQQYKFLPTEL REDEG 3441 Ā 3 1584 NSARGGVGVRGARAMATVQEKAAALNLSALHS PAHRPPGFSVAQKPFGATYVWSSIINTLQTQVEV KKRRHRLKRHNDCFVGSEAVDVIFSHLIQNKYF **GDVDIPRAKVVRVCQALMDYKVFEAVPTKVFG** KDKKPTFEDSSCSLYRFTTIPNQDSQLGKENKLY SPARYADALFKSSDIRSASLEDLWENLSLKPANS PHVNISTTLSPQVINEVWQEETIGRLLQLVDLPLL DSLLKQQEAVPKIPQPKRQSTMVNSSNYLDRGIL KAYSDSQEDEWLSAAIDCLEYLPDQMVVEISRSF PEQPDRTDLVKELLFDAIGRYYSSREPLLNHLSD VHNGIAELLVNGKTEIALEATQLLLKLLDFQNRE **EFRRLLYFMAVAANPSEFKLQKESDNRMVVKRI** FSKAIVDNKNLSKGKTDLLVLFL\MDHQKDVFKI PGTL\HKIVS\VK\LMAIQNGRDPNRDAGYIYCQRI DQRDYSNITEKTTIDELLYLLKTLDEDSKLSAKE KKK\LLGQFYKCHPDIFIEHFGD 3442 160 822 Â SPASGHCRLNGAAVAMFGCLVAGRLVQTAAQQ VAEDKFVFDLPDYESINHVVVFMLGTIPFPEGMG GSVYFSYPDSNGMPVWQLLGFVTNGKPSAIFKIS GLKSGEGSQHPFGAMNIVRTPSVAQIGISVELLDS MAQQTPVGNAAVSSVDSFTQFTQKMLDNFYNF ASSFAVSQ/VPDDTQ/RPSEMFIPANVVLKWYENF **QRRTSTEPSLLENIIWIKINF** <u>3443</u> 1373 A **SWHVRRRWLEATMAGGMKVAVSPAVGPGPWG** SGVGGGGTVRLLLILSGCLVYGTAETDVNVVML QESQVCEKRASQQFCYTNVLIPQWHDIWTRIQIR VNSSRLVRVTQVENEEKLKELEQFSIWNFFSSFL KEKLNDTYVNVGLYSTKTCLKVEIIEKDTKYSVI VIRRFDPKLFLVFLLGLMLFFCGDLLSRSQIFYYS TGMTVGIVASL\LIIIFILSKFMPKKSPIYVILVGGW SFSLYLIQLVFKNLQEIWRCYWQYLLSYVLTVGF MSFAVCYKYGPLENERSINLLTWTLOLMGLCFM YSGIQIPHIALAIIIIALCTKNLEHPIQWLYITCRKV CKGAEKPVPPRLLTEEEYRIQGEVETRKALEELR **EFCNSPDCSAWKTVSRIQSPKRFADFVEGSSHLT PNEVSVHEQEYGLGSIIAQDEIYEEASSEEEDSYS** RCPAITONNFLT 3444 566 1718 A KGLERTCCAMEESDSEKTTEKENLGPRMDPPLG EPG\GSLGWVLPNTAMKKKVLLMGKSGSGKTS

> MRSIIFANYIARDTRRLGATILDRIHSLQINSSLST YSLVDSVGNTKTFDVEHSHVRFLGNLVLNLWDC GGQDTFMENYFTSQRDNIFRNVEVLIYVFDVESR ELEKDMHYYQSCLEAILQNSPDAKIFCLVHKMD LVQEDQRDLIFKEREEDLRRLSRPLECSCFRTSIW

SEQ ID NO:	Method	Predicted beginning nucleotide location corresponding to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion
				DETLYKAWSSIVYQLIPNVQQLEMNLRNFAEIIE ADEVLLFERATFLVISHYQCKEQRDAHRFEKISNI IKQFKLSCSKLAASFQSMEVRNSNFAAFIDIFTSN TYVMVVMSDPSIPSAATLINIRNARKHFEKLERV DGPKQCLLMR
3445	A	566	1718	KGLERTCCAMEESDSEKTTEKENLGPRMDPPLG EPG\GSLGWVLPNTAMKKKVLLMGKSGSGKTS MRSIIFANYIARDTRRLGATILDRIHSLQINSSLST YSLVDSVGNTKTFDVEHSHVRFLGNLVLNLWDC GGQDTFMENYFTSQRDNIFRNVEVLIYVFDVESR ELEKDMHYYQSCLEAILQNSPDAKIFCLVHKMD LVQEDQRDLIFKEREEDLRRLSRPLECSCFRTSIW DETLYKAWSSIVYQLIPNVQQLEMNLRNFAEIIE ADEVLLFERATFLVISHYQCKEQRDAHRFEKISNI IKQFKLSCSKLAASFQSMEVRNSNFAAFIDIFTSN TYVMVVMSDPSIPSAATLINIRNARKHFEKLERV DGPKQCLLMR
3446	A	566	1718	KGLERTCCAMEESDSEKTTEKENLGPRMDPPLG EPG\GSLGWVLPNTAMKKKVLLMGKSGSGKTS MRSIIFANYIARDTRRLGATILDRIHSLQINSSLST YSLVDSVGNTKTFDVEHSHVRFLGNLVLNLWDC GGQDTFMENYFTSQRDNIFRNVEVLIYVFDVESR ELEKDMHYYQSCLEAILQNSPDAKIFCLVHKMD LVQEDQRDLIFKEREEDLRRLSRPLECSCFRTSIW DETLYKAWSSIVYQLIPNVQQLEMNLRNFAEIIE ADEVLLFERATFLVISHYQCKEQRDAHRFEKISNI IKQFKLSCSKLAASFQSMEVRNSNFAAFIDIFTSN TYVMVVMSDPSIPSAATLINIRNARKHFEKLERV DGPKQCLLMR
3447	A		2930	VLLGPLWDKLSTADHPVIVTMASKRKSTTPCMIP VKTVVLQDASMEAQPAETLPEGPQQDLPPEASA ASSEAAQNPSSTDGSTLANGHRSTLDGYLYSCK YCDFRSHDMTQFVGHMNSEHTDFNKDPTFVCSG CSFLAKTPEGLSLHNATCHSGEASFVWNVAKPD NHVVVEQSIPESTSTPDLAGEPSAEGADGQAEIIIT KTPIMKIMKGKAEAKKIHTLKENVPSQPVGEALP KLSTGEMEVREGDHSFINGAVPVRQASASSAKN PHAANGPLIGTVPVLPAGIAQFLSLQQQPPVHAQ HHVHQPLPTAKALPKVMIPLSSIPTYSAAMDSNS FLKNSFHKFPYPTKAELCYLTVVTKYPEEQLKIW FTAQRLKQGISWSPEEIEDARKKMFNTVIQSVPQ PTITVLNTPLVASAGNVQHLIQAALPGHVVGQPE GTGGGLLVTQPLMANGLQATSSPLPLTVTSVPK QPGVAPINTVCSNTTSAVKVVNAAQSLLTACPSI TSQAFLDASIYKNKKSHEQLSALKGSFCRNQFPG QSEVEHLTKVTGLSTREVRKWFSDRRYHCRNLK GSRAMIPGDHRSIIIDSVPEVSFSPSSKVPEVTCIPT TATLATHPSAKRQSWHQTPDFTPTKYKERAPEQ LRALESSFAQNPLPLDEELDRLRSETKMTRREIDS WFSERRKKVNAEETKKAEENASQEEEEAAEDEG GEEDLASELRVSGENGSLEMPSSHILAERKVSPIK INLKNLRVTEANGRNEIPGLGACDPEDDESNKLA EQLPGKVSCKKTAQQRHLLRQLFVQTQWPSNQD YDSIMAQTGLPRPEVVRWFGDSRYALKNGQLK WYEDYKRGNFPPGLLVIAPGNRELLODYYMTHK

SEQ ID NO:	Method	Predicted beginning nucleotide location corresponding to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Prollne, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \text{\tex{\tex
				MLYEEDLQNLCDKTQMSSQQVKQWFAEKMGEE TRAVADTGSEDQGPGTGELTAVHKGMGDTYSE VSENSESWEPRVPEASSEPFD\TSSPQAGRQLETD
3448	A			FVARAEKGFRTREAHLLQVAGVGTGLQNGASLS GLASGVMAQRAFPNPYADYNKSLAEGYFDAAG RLTPEFSQRLTNKIRELLQQMERGLKSADPRDGT GYTGWAGIAVLYLHLYDVFGDPAYLQLAHGYV KQSLNCLTKRSITFLCGDAGPLAVAAVLYHKMN NEKQAEDCITRLIHLNKIDPHAPNEMLYGRIGYIY ALLFVNKNFGVEKIPQSHIQQICETILTSGENLAR KRNFTAKSPLMYEWYQEYYVGAAHGLAGIYYY LMQPSLQVSQGKLHSLVKPSVDYVCQLKFPSGN YPPCIGDNRDLLVHWCHGAPGVIYMLIQAYKVF R/EREKYLC\DAYQCADVIWQYGLLKKGYGLCY\ GSAGNAYAFLTLYNLTQDMKYLYRACKFAEWC LEYGEHGCRTPDTPFSLFEGMAGTIYFL\ADLLFP TKAR\FPAFEL
3449	A	3	2389	SRHVTGAARSPSRAGPSDPPAMGDEDDDESCAV ELRITEANLTGHEEKVSVENFELLKVLGTGAYGK VFLVRKAGGHDAGKLYAMKVLRKAALVQRAK TQEHTRTERSVLELVRQAPFLVTLHYAFQTDAKL HLILDYVSGGEMFTHLYQRQYFKEAEVRVYGGE IVLALEHLHKLGIIYRDLKLENVLLDSEGHIVLTD FGLSKEFLTEEKERTFSFCGTIEYMAPEIIRSKTGH GKAVDWWSLGILLFELLTGASPFTLEGERNTQAE VSRRILKCSPPFPPRIGPVAQDLLQRLLCKDPKKR LGAGPQGAQEVRNHPFFQGLDWVALAARKIPAP FRPQIRSELDVG\NFAEEFTRLEPVYSPPGQ\PPPG DPRIFQGYSFVAPSILFDHNNAVMTDGLEAPGAG DRPGRAAVARSAMMQDSPFFQQYELDLREPALG QGSFSVCRRCRQRQSGQEFAVKILSRRLEANTQR EVAALRLCQSHPNVVNLHEVHHDQLHTYLVLEL LRGGELLEHIRKKRHFSESEASQILRSLVSAVSFM HEEAGVVHRDLKPENILYADDTPGAPVKIIDFG/F SPRLRPQSPGVPMQTPSFTLQYAAPELLAQQGYD ESCDLWSLGVILY\MMLSGQAPFQGASGQGGQS QAAEIMCKIREGRFSLDGEAWQGVSEEAKELVR GLLTVDPAKRLKLEGLRGSSWLQDGSARSSPPLR TPDVLESSGPAVRSGLNATFMAFNRGKREGFFLK SVENAPLAKRRKQKLRSATASRRGSPAPANPGR APVASKGAPRANGPLPPS
3450	A	201	1705	KGTEMNKSRWQSRRRHGRRSHQQNPWFRLRDS EDRSDSRAAQPAHDSGHGDDESPSTSSGTAGTSS VPELPGFYFDPEKKRYFRLLPGHNNCNPLTKESIR QKEMESKRLRLLQEEDRRKKIARMGFNASSMLR KSQLGFLNVTNYCHLAHELRLSCMERKKVQIRS MDPSALASDRFNLILADTNSDRLFTVNDVTVGGS KYGIINLQSLKTPTLKVFMHENLYFTNRKV\NSV CWASLNHLDSHILLCLMGLAETPGCATLLPASLF VNSHPAGIDRPG\MLCSFRIPGAWSCAWSLNIQA NNCFSTGLSRRVLLTNVVTGHRQSFGTNSDVLA QQFALMAPLLFNGCRSGEIFAIDLRCGNQGKGW KATRLFHDSAVTSVRILQDEQYLMASDMAGKIK LWDLRTTKCVRQYEGHVNEYAYLPLHVHEEEGI LVAVGQDCYTRIWSLHDARLLRTIPSPYPASKAD

محددتات.

SEQ ID	Method	Predicted	Predicted end	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid,
NO:		beginning	nucleotide	E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine,
		nucleotide location	location corresponding	N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine,
		corresponding	to last amino	T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine,
		to first amino	acid residue of	X=Unknown, *=Stop codon, /=possible nucleotide deletion,
		acid residue of peptide	peptide sequence	=possible nucleotide insertion
		sequence	<u>-</u>	IDONA POODY COOR CAROLLY MANGODI WOYOYO
2451	<u> </u>	19	6033	IPSVAFSSRLGGSRGAPGLLMAVGQDLYCYSYS LLSAMLSHGAGLALWITLSLLQTGLAEPERCNFT
3451	A		0033	LAESKASSHSVSIQWRILGSPCNFSLIYSSDTLGA
				ALCPTFRIDNTTYGCNLQDLQAGTIYNFKIISLDE
				ERTVVLQTDPLPPARFGVSKEKTTSTGLHVWWT
				PSSGKVTSYEVQLFDENNQKIQGVQIQESTSWNE
				YTFFNLTAGSKYNIAITAVSGGKRSFSVYTNGST
				VPSPVKDIGISTKANSLLISWSHGSGNVERYRLM
				LMDKGILVHGGVVDKHATSYAFHGLSPGYLYNL
				TVMTEAAGLQNYRWKLVRTAPMEVSNLKVTND
				GSLTSLKVKWQRPPG\NVDSYNITLSHKGTIKESR
				VLAPWIT\ETHFKELVPGRLY\QVTCSAVSLGELS
				AQKM\AVGRTFPDKVANLEANNNGRMRSLVVS
				WSPPAGDWEQYRILLFNDSVVLLNITVGKEETQ YVMDGTGLVPGRQYEVEVIVESGNLKNSERCQG
				RTVPLAVLQLRVKHANETSLSIMWQTPVAEWEK
				YIISLADRDLLLIHKSLSKDAKEFTFTDLVPGRKY
				MATVTSISGDLKNSSSVKGRTVPAQVTDLHVAN
				QGMTSSLFTNWTQAQGDVEFYQVLLIHENVVIK
ļ				NESISSETSRYSFHSLKSGSLYSVVVTTVSGGISSR
				QVVVEGRTVPSSVSGVTVNNSGRNDYLSVSWLL
				APGDVDNYEVTLSHDGKVVQSLVIAKSVRECSF
				SSLTPGRLYTVTITTRSGKYENHSFSQERTVPDKV
				QGVSVSNSARSDYLRVSWVHATGDFDHYEVTIK
				NKNNFIQTKSIPKSENECVFVQLVPGRLYSVTVT
<u> </u>			·	TKSGQYEANEQGNGRTIPEPVKDLTLRNRSTEDL HVTWSGANGDVDQYEIQLLFNDMKVFPPFHLVN
:				TATEYRFTSLTPGRQYKILVLTISGDVQQSAFIEG
				FTVPSAVKNIHISPNGATDSLTVNWTPGGGDVDS
				YTVSAFRHSQKVDSQTIPKHVFEHTFHRLEAGEQ
				YQIMIASVSGSLKNQINVVGRTVPASVQGVIADN
				AYSSYSLIVSWQKAAGVAERYDILLLTENGILLR
				NTSEPATTKQHKFEDLTPGKKYKIQILTVSGGLFS
				KEAQTEGRTVPAAVTDLRITENSTRHLSFRWTAS
				EGELSWYNIFLYNPDGNLQERAQVDPLVQSFSFQ
				NLLQGRMYKMVIVTHSGELSNESFIFGRTVPASV SHLRGSNRNTTDSLWFNWSPASGDFDFYELILYN
				PNGTKKENWKDKDLTEWRFQGLVPGRKYVLW
				VVTHSGDLSNKVTAESRTAPSPPSLMSFADIANT
				SLAITWKGPPDWTDYNDFELQWLPRDALTVFNP
				YNNRKSEGRIVYGLRPGRSYQFNVKTVSGDSWK
				TYSKPIFGSVRTKPDKIQNLHCRPQNSTAIACSWI
				PPDSDFDGYSIECRKMDTQEVEFSRKLEKEKSLL
				NIMMLVPHKRYLVSIKVQSAGMTSEVVEDSTIT
				MIDRPPPPPHIRVNEKDVLISKSSINFTVNCSWFS
				DTNGAVKYFTVVVREADGSDELKPEQQHPLPSY
				LEYRHNASIRVYQTNYFASKCAENPNSNSKSFNI
				KLGAEMESLGGKCDPTQQKFCDGPLKPHTAYRI SIRAFTQLFDEDLKEFTKPLYSDTFFSLPITTESEP
				LFGAIEGVSAGLFLIGMLVAVVALLICROKVSHG
				RERPSARLSIRRDRPLSVHLNLGQKGNRKTSCPIK
				INQFEGHFMKLQADSNYLLSKEYEELKDVGRNQ
				SCDIALLPENRGKNRYNNILPYDATRVKLSNVDD
				DPCSDYINASYIPGNNFRREYIVTQGPLPGTKDDF
				WKMVWEQNVHNIVMVTQCVEKGRVKCDHYW

SEQ ID NO:	Method	Predicted beginning nucleotide location corresponding to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion
				PADQDSLYYGDLILQMLSESVLPEWTIREFKICGE EQLDAHRLIRHFHYTVWPDHGVPETTQSLIQFVR TVRDYINRSPGAGPTVVHCSAGVGRTGTFIALDR ILQQLDSKDSVDIYGAV\HDLRLHRVHMVQTEC QYVYLHQCVRDVLRARKLRSEQENPLFPIYENV NPEYHRDPVYSRH
3452	A	63	1073	FFRSSSDNGSPIRQYE/HSTPAHQGPVMGLEGKS/ARNSQLRIVLVGKTGAGKSATGNSILGRKVFHSGTAAKSITKKCEKRSSSWKETELVVVDTPGIFDTEVPNAETSKEIIRCILLTSPGPHALLLVVPLGRYTEEEHKATEKILKMFGERARSFMILIFTRKDDLGDTNLHDYLREAPEDIQDLMDIFGDRYCALNNKATGAEQEAQRAQLLGLIQRVVRENKEGCYTNRMYQRAEEEIQKQTQAMQELHRVELEREKARIREEYEEKIRKLEDKVEQEKRKKQMEKKLAEQEAHYAVRQQRARTEVESKDGILELIMTALQIASFILLRLFAED
3453	A	2674	514	GPITFLKKKAKMKDMPLRIHVLLGLAITTLVQAV DKKVDCPRLCTCEIRPWFTPRSIYMEASTVDCND LGLLTFPARLPANTQILLLQTNNIAKIEYSTDFPV NLTGLDLSQNNLSSVTNINGKKMPQLLSVYLEEN KLTELPEKCLSELSNLQELYINHNLLSTISPGAFIG LHNLLRLHLNSNRLQMINSKWFDALPNLEILMIG ENPIIRIKDMNFKPLINLRSLVIAGINLTEIPDNAL VGLENLESISFYDNRLIKVPHVALQKVVNLKFLD LNKNPINRIRRGDFSNMLHLKELGINNMPELISID SLAVDNLPDLRKIEATNNPRLSYIHPNAFFRLPKL ESLMLNSNALSALYHGTIESLPNLKEISIHSNPIRC DCVIRWMNMNKTNIRFMEPDSLFCVDPPEFQGQ NVRQVHFRDMMEICLPLIAPESFPSNLNVEAGSY VSFHCRATA\EPQPEIYWITPSGQKLLPNT\LTDKF YVHSEGTLDINGVTPKEGGLYTCIATNLVGADLK SVMIKVDGSFPQDNNGSLNIKIRDIQANSVLVSW KASSKILKSSVKWTAFVKTENSHAAQSARIPSDV KVYNLTHLNPSTEYKICIDIPTIYQKNRKKCVNVT TKGLHPDQKEYEKNNTTTLMACLGGLLGIIGVIC LISCLSPEMNCDGGHSYVRNYLQKPTFALGELYP PLINLWEAGKEKSTSLKVKATVIGLPTNMS
3454	A	1844	244	ERYLFATYVAPSATLDIGLQQEKKKEIYMKIQPP FEDLFDTAEEYILLLLEPWTKMVKSDQIAYKKV ELVEETRQLDSTYFRKLQALHKETFSKKAEDTTC EIGTGILSLSNVSKRTEYWDNVPAEYKHFKFSDL LNNKLEFEHFRQFLETHSSSMDLMCWTDIEQFRR ITYRDRNQRKAKSIYIKNKYLNKKYFFGPNSPAS LYQQNQVMHLSGGWGKILHEQLDAPVLVEIQK HVQNRLENVWLPLFLASEQFAARQKIKVQMKDI AEELLLQKAEKKIGVWKPVESKWISSSCKIIAFRK ALLNPVTSRQFQRFVALKGDLLENGLLFWQEVQ KYKDLCHSHCDESVIQKKITTIINCFINSSIPPALQI DIPVEQAQKIIEHRKELGPYVFREAQMTFLGVMF KFWPQFCEFRKNLTDENIMSVLERRQEYNKQKK KLAVL/QNDEKSGKDGIKQYANTSVPAIKTALLS DSFLGLQPYGRQPTWCYSKYIEALEQERILLKIQE ELEK\SCLQACNLSQILRLALQLCL
3455	A	228	3330	APTAQAMMSFGGADALLGAPFAPLHGGGSLHY ALARKGGAGGTRSAAGSSSGFHSWTRTSVSSVS

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SEQ ID No: Method Predicted beginning nucleotide location corresponding to first amino acid residue of peptide sequence Predicted end sequence Predicted end location corresponding to first amino acid residue of peptide sequence Predicted end sequence (sed end end end end end end end end end e
nucleotide location corresponding corresponding to first amino acid residue of peptide sequence Sequence Incation corresponding to first amino acid residue of peptide sequence Incation corresponding to first amino acid residue of peptide sequence Incation corresponding to first amino acid residue of peptide sequence Incation corresponding to first amino acid residue of peptide sequence Incation corresponding to first amino acid residue of peptide sequence Incation corresponding to first amino acid residue of peptide sequence Incation corresponding to first amino acid residue of peptide sequence Incation corresponding to first amino acid residue of peptide sequence Incation corresponding to first amino acid residue of peptide sequence Incation corresponding to first amino acid residue of peptide sequence Incation corresponding corresp
corresponding to first amino acid residue of peptide sequence Table
acid residue of peptide sequence S=Unknown, *=Stop codon, /=possible nucleotide deletion
acid residue of peptide sequence ASPSRFRGAGAASSTDSLDTLSNGPEGCMV. TSRSEKEQLQALNDRFAGYIDKVRQLEAHN GEAAALRQQAGRSAMGELYEREVREMRC. RLGAARGQLRLEQEHLLEDIAHVRQRLDDE REEAEAAARALARFAQEAEAARVDLQKKA QEECGYLRRHHQEEVGELLGQIQGSGAAQA QAETRDALKCDVTSALREIRAQLEGHAVQS SEEWFRVRLDRLSEAAKVNTDAMRSAQEEL RRQLQARTTELEALKSTKDSLERQRSELEDI DIASYQEAIQQLDAELRNTK WEMAAQLREV LNVKMALDIEIAAYRKLLEGEECRIGFGPIPHEGLPKIPSVSTHIKVKSEEKIKVVEKSEKETV QTEETQVTEEVTEEDKEAKEEGKEEGG AEGGEETKSPPAEEAASPEKEAKSPVKEEA AEAKSPEKEAKSPVKEEA AEAKSPEKAESPEKAKSPAKSPAKSPEKAKSPKEE SPAEVKSPEKAKSPAKSPEKAKSPKEE SPAEVKSPEKAKSPKEEAKSPEKAKSPKEEKEERKSPEKAKSPVKEEAKSPKEEKEEKEEARPATPKTEJ DSKKEEAPKKEAPKKEPKPKEEKEEVKEEVKEEAKSPKKEEARPAKEAPAKVEVKEDAEKEEARAPAKEEARAPAKVEVKEDAEKEEAKSPAKEEARAPKEEPKKEEVKEEAKSPAKEEARAPAKVEVKEDAEKEEAKSPKEAEKEEARAPAKVEVKEDAEKEEAKSPKEEKEEARAPAKEEVKEEAKEEARAPAKEEARAPAKEEVKEEAKEEARAPAKEEARAPAKEEARAPAKEEARAPAKEEARARAEEARARAEEARARAEAREEARARAEAREEARARAEAREEARARAEAREEARARAEAREEARARAEAREEARARAEAREEARARAEAREEARARAEAREEARARAEAREE
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ASPSRFRGAGAASSTDSLDTLSNGPEGCMV TSRSEKEQLQALNDRFAGYIDKVRQLEAHN GEAAALRQQAGRSAMGELYEREVREMRC RLGAARGQLRLEQEHLLEDIAHVRQRLDDE REEAEAAARALARFAQEAEAARVDLQKKA QEECGYLRRHHQEEVGELLGQIQGSGAAQA QAETRDALKCDVTSALREIRAQLEGHAVQS SEEWFRVRLDRLSEAAKVNTDAMRSAQEE! RRQLQARTTELEALKSTKDSLERQRSELEDI DIASYQEAIQQLDAELRNTKWEMAAQLREY LNVKMALDIEIAAYRKLLEGEECRIGFGPIPH EGLPKIPSVSTHIKVKSEEKIKVVEKSEEKITV QTEETQVTEEVTEEDKEAKEEGKEEEGG AEGGEETKSPPAEEAASPEKEAKSPVKEEA AEAKSPEKEEAKSPAEVKSPEKAKSPVKEEA SPAEAKSPEKAKSPVKEEAKSPEKAKSPVKE SPAEVKSPEKAKSPVKEEAKSPEKAKSPVKE EKEEAKSPEKAKSPVKEEAKSPEKAKSPVKE KSPEKAKSPVKEEAKSPVKEEAKSPKE EKEEAKSPEKAKSPVKEEAKSPVKEEAKSP EKEEAKSPEKAKSPVKEEAKSPKEEAKSP EKEEAKSPEKAKSPVKEEAKSPEKAKSPVK KSPEKAKSPVKEEAKSPEKAKSPVKEEAKSP EKEEAKSPEKAKSPVKEEAKSPEKAKSPVK EKEEAKSPEKAKSPVKEEAKSPEKAKSPVK EKSPEKAKSPVKEEAKSPEKAKSPVKEEAKSP EKEEAKSPEKAKSPVKEEAKSPEKAKSPVK ESPVKEEAKTPEKAKSPVKEEAKSPEKAKSP KTLDVKSPEAKTPAKEEARSPADKFPEKAK EEVKSPEKAKSPLKEDAKAPEKEIPKEEVK KEEKPQEVKVKEPPKKAEEEKAPATPKTE DSKKEEAPKKEAPKPKVEEKKEPAVEKPKE EAKKEEAEDKKKVPTPEKEAPAAVEVYKEDA EKTEVAKKEPDDAKAKEPSKPAEKKEAAPE
TSRSEKEQLQALNDRFAGYIDKVRQLEAHN GEAAALRQQQAGRSAMGELYEREVREMRC RLGAARGQLRLEQEHLLEDIAHVRQRLDDE REEAEAAARALARFAQEAEAARVDLQKKA QEECGYLRRHHQEEVGELLGQIQGSGAAQA QAETRDALKCDVTSALREIRAQLEGHAVQS SEEWFRVRLDRLSEAAKVNTDAMRSAQEEI RRQLQARTTELEALKSTKDSLERQRSELEDI DIASYQEAIQQLDAELRNTKWEMAAQLREY LNVKMALDIEIAAYRKLLEGEECRIGFGPIPF EGLPKIPSVSTHIKVKSEEKIKVVEKSEKETV QTEETQVTEEVTEEDKEAKSEEGKEEEGG AEGGEETKSPPAEEAASPEKAKSPAKEEA AEAKSPEKEEAKSPAEVKSPEKAKSPAKEEA SPAEAKSPEKAGSPVKEEAKSPAKESPAKE SPAEVKSPEKAKSPVKEEAKSPEKAKSPVKEEA EKEEAKSPEKAKSPVKEEAKSPEKAKSPVK KSPEKAKSPVKEEAKSPEKAKSPVK KSPEKAKSPVKEEAKSPEKAKSPVK KSPEKAKSPVKEEAKSPEKAKSPVK KSPEKAKSPVKEEAKSPEKAKSPVK KSPEKAKSPVKEEAKSPEKAKSPVK KSPEKAKSPVKEEAKSPEKAKSPVK KSPEKAKSPVKEEAKSPEKAKSPVK KSPEKAKSPVKEEAKSPEKAKSPVK KSPEKAKSPVKEEAKSPEKAKSPVK KSPEKAKSPVKEEAKSPEKAKSPVK KSPEKAKSPVKEEAKSPEKAKSPVK KSPEKAKSPVKEEAKSPEKAKSP KTLDVKSPEAKTPAKEEARSPADKFPEKAK EEVKSPEKAKSPLKEDAKAPEKEPKKEEVK KEEEKPQEVKVKEPPKKAEEEKAPATPKTEI DSKKEEAPKKEAPKPKVEEKKEPAVEKPKE EAKKEEAEDKKKVPTPEKEAPAKVEVKEDA EKTEVAKKEPDDAKAKEPSKPAEKKEAAPE
GEAAALRQQAGRSAMGELYEREVREMRO RLGAARGQLRLEQEHLLEDIAHVRQRLDDE REEAEAAARALARFAQEAEAARVDLQKKA QEECGYLRRHHQEEVGELLGQIQGSGAAQA QAETRDALKCDVTSALREIRAQLEGHAVQS SEEWFRVRLDRLSEAAKVNTDAMRSAQEEI RRQLQARTTELEALKSTKDSLERQRSELEDI DIASYQEAIQQLDAELRNTKWEMAAQLREY LNVKMALDIEIAAYRKLLEGEECRIGFGPIPE EGLPKIPSVSTHIKVKSEEKIKVVEKSEKETV QTEETQVTEEVTEEDKEAKEEGKEEEGG AEGGEETKSPPAEEAASPEKAKSPVKEEA AEAKSPEKEEAKSPAEVKSPEKAKSPAKEA SPAEAKSPEKAGNFQAEVKSPEKAKSPAKEA SPAEAKSPEKAKSPVKEEAKSPEKAKSPKE SPAEVKSPEKAKSPVKEEAKSPEKAKSPVK SPAEVKSPEKAKSPVKEEAKSPEKAKSPVK KSPEKAKSPVKEEAKSPEKAKSPVK KSPEKAKSPVKEEAKSPEKAKSPVK KSPEKAKSPVKEEAKSPEKAKSPVK KSPEKAKSPVKEEAKSPEKAKSP KSPVKEEAKTPEKAKSPVKEEAKSP KTLDVKSPEAKTPAKAEARSPADKFPEKAK EEVKSPEKAKSPLKEDAKAPEKEEVK KEEEKPQEVKVKEPPKKAEEEKAPATPKTE DSKKEEAPKKEAPKPKVEEKKEPAVEKPED EKTEVAKKEPDDAKAKEPSKPAEKKEAPE
RLGAARGQLRLEQEHLLEDIAHVRQRLDDE REEAEAAARALARFAQEAEAARVDLQKKA QEECGYLRRHHQEEVGELLGQIQGSGAAQA QAETRDALKCDVTSALREIRAQLEGHAVQS SEEWFRVRLDRLSEAAKVNTDAMRSAQEE RRQLQARTTELEALKSTKDSLERQRSELEDI DIASYQEAIQQLDAELRNTKWEMAAQLREY LNVKMALDIEIAAYRKLLEGEECRIGFGPIPH EGLPKIPSVSTHIKVKSEEKIKVVEKSEKETV QTEETQVTEEVTEEEDKEAKEEEGKEEEGG AEGGEETKSPPAEEAASPEKEAKSPVKEEA AEAKSPEKEEAKSPAEVKSPEKAKSPVKEEA SPAEAKSPEKOGKQNFQAEVKSPEKAKSPVKE SPAEVKSPEKAKSPVKEEAKSPEKAKSPVKE SPAEVKSPEKAKSPVKEEAKSPEKAKSPVKE EKEEAKSPEKAKSPVKEEAKSPEKAKSPVK KSPEKAKSPVKEEAKSPEKAKSPVK KSPEKAKSPVKEEAKSPEKAKSPVK KSPVKEEAKTPEKAKSPVKEEAKSPEKAKSI KTLDVKSPEAKTPAKEEARSPADKFPEKAK EEVKSPEKAKSPLKEDAKAPEKEIPKKEEVK KEEERPQEVKVKEPPKKAEEKAPATPKTE DSKKEEAPKKEAPKVEEKKEPAKVEVKEDA EKTEVAKKEPDDAKAKEPSKPAEKKEAAPE
REEAEAAARALARFAQEAEAARVDLQKKA QEECGYLRRHHQEEVGELLGQIQGSGAAQA QAETRDALKCDVTSALREIRAQLEGHAVQS SEEWFRVRLDRLSEAAKVNTDAMRSAQEEI RRQLQARTTELEALKSTKDSLERQRSELEDI DIASYQEAIQQLDAELRNTKWEMAAQLREY LNVKMALDIEIAAYRKLLEGEECRIGFGPIPH EGLPKIPSVSTHIKVKSEEKIKVVEKSEKETV QTEETQVTEEVTEEDKEAKBEEGKEEGGG AEGGEETKSPPAEEAASPEKEAKSPVKEEA AEAKSPEKEEAKSPAEVKSPEKAKSPAKEEA PE\AKSPEKDGKQNFQAEVKSPEKAKSPAKE SPAEAKSPEKAKSPVKEEAKSPEKAKSPVKE SPAEVKSPEKAKSPVKEEAKSPEKAKSPVKE EKEEAKSPEKAKSPVKAEAKSPEKAKSPVK KSPEKAKSPVKEEAKSPEKAKSPVKEEAKSP KSPVKEEAKTPAKEEARSPADKFPEKAKS KTLDVKSPEAKTPAKEEARSPADKFPEKAK EEVKSPEKAKSPLKEDAKAPEKEIPKKEEVK KEEEKPQEVKVKEPPKKAEEEKAPATPKTE DSKKEEAPKKEAPKPKVEEKAPAVEVKEDA EKTEVAKKEPDDAKAKEPSKPAEKKEAAPE
QEECGYLRRHHQEEVGELLGQIQGSGAAQA QAETRDALKCDVTSALREIRAQLEGHAVQS SEEWFRVRLDRLSEAAKVNTDAMRSAQEEI RRQLQARTTELEALKSTKDSLERQRSELEDH DIASYQEAIQQLDAELRNTKWEMAAQLREY LNVKMALDIEIAAYRKLLEGEECRIGFGPIPH EGLPKIPSVSTHIKVKSEEKIKVVEKSEKETV QTEETQVTEEVTEEEDKEAKEEGKEEEGG AEGGEETKSPPAEEAASPEKEAKSPVKEEA AEAKSPEKEEAKSPAEVKSPEKAKSPAKEE PE\AKSPEKDGKQNFQAEVKSPEKAKSPAKE SPAEAKSPEKAKSPVKEEAKSPEKAKSPAKE SPAEAKSPEKAKSPVKEEAKSPEKAKSPEKA SPAEVKSPEKAKSPVKAEAKSPEKAKSPVK EKEEAKSPEKAKSPVKEEAKSPEKAKSPVK KSPEKAKSPVKEEAKSPEKAKSPVK KSPEKAKSPVKEEAKSPEKAKSPVK KSPEKAKSPVKEEAKSPVKEEAKSP KTLDVKSPEAKTPAKEEARSPADKFPEKAK EEVKSPEKAKSPLKEDAKAPEKEIPKKEEVK KEEEKPQEVKVKEPPKKAEEEKAPATPKTEI DSKKEEAPKKEAPKPKVEEKKEPAVEKPKE EAKKEEAEDKKKVPTPEKEAPAKVEVKEDA EKTEVAKKEPDDAKAKEPSKPAEKKEAAPE
QEECGYLRRHHQEEVGELLGQIQGSGAAQA QAETRDALKCDVTSALREIRAQLEGHAVQS SEEWFRVRLDRLSEAAKVNTDAMRSAQEEI RRQLQARTTELEALKSTKDSLERQRSELEDH DIASYQEAIQQLDAELRNTKWEMAAQLREY LNVKMALDIEIAAYRKLLEGEECRIGFGPIPH EGLPKIPSVSTHIKVKSEEKIKVVEKSEKETV QTEETQVTEEVTEEEDKEAKEEGKEEEGG AEGGEETKSPPAEEAASPEKEAKSPVKEEA AEAKSPEKEEAKSPAEVKSPEKAKSPAKEE PE\AKSPEKDGKQNFQAEVKSPEKAKSPAKE SPAEAKSPEKAKSPVKEEAKSPEKAKSPAKE SPAEAKSPEKAKSPVKEEAKSPEKAKSPEKA SPAEVKSPEKAKSPVKAEAKSPEKAKSPVK EKEEAKSPEKAKSPVKEEAKSPEKAKSPVK KSPEKAKSPVKEEAKSPEKAKSPVK KSPEKAKSPVKEEAKSPEKAKSPVK KSPEKAKSPVKEEAKSPVKEEAKSP KTLDVKSPEAKTPAKEEARSPADKFPEKAK EEVKSPEKAKSPLKEDAKAPEKEIPKKEEVK KEEEKPQEVKVKEPPKKAEEEKAPATPKTEI DSKKEEAPKKEAPKPKVEEKKEPAVEKPKE EAKKEEAEDKKKVPTPEKEAPAKVEVKEDA EKTEVAKKEPDDAKAKEPSKPAEKKEAAPE
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SEEWFRVRLDRLSEAAKVNTDAMRSAQEEJ RRQLQARTTELEALKSTKDSLERQRSELEDI DIASYQEAIQQLDAELRNTKWEMAAQLREY LNVKMALDIEIAAYRKLLEGEECRIGFGPIPE EGLPKIPSVSTHIKVKSEEKIKVVEKSEKETV QTEETQVTEEVTEEDKEAKEEGKEEEGG AEGGEEETKSPPAEEAASPEKEAKSPVKEEA AEAKSPEKEEAKSPAEVKSPEKAKSPAKEE PE\AKSPEKDGKQNFQAEVKSPEKAKSPAKE SPAEAKSPEKAKSPVKEEAKSPEKAKSPVKE SPAEVKSPEKAKSPVKAEAKSPEKAKSPVK EKEEAKSPEKAKSPVKAEAKSPVKEEAKSP KSPVKEEAKSPVKEEAKSPVKEEAKSP KSPVKEEAKTPEKAKSPVKEEAKSPEKAKSP KTLDVKSPEAKTPAKEEARSPADKFPEKAKSEVK EEVKSPEKAKSPLKEDAKAPEKEIPKKEEVK KEEEKPQEVKVKEPPKKAEEEKAPATPKTEI DSKKEEAPKKEAPKVEEKEPAVEKPKE EAKKEEAEDKKKVPTPEKEAPAKVEVKEDA EKTEVAKKEPDDAKAKEPSKPAEKKEAAPE
RRQLQARTTELEALKSTKDSLERQRSELEDI DIASYQEAIQQLDAELRNTKWEMAAQLREY LNVKMALDIEIAAYRKLLEGEECRIGFGPIPE EGLPKIPSVSTHIKVKSEEKIKVVEKSEKETV QTEETQVTEEVTEEEDKEAKEEEGKEEEGG AEGGEEETKSPPAEEAASPEKEAKSPVKEEA AEAKSPEKEEAKSPAEVKSPEKAKSPAKEEA PE\AKSPEKDGKQNFQAEVKSPEKAKSPAKE SPAEAKSPEKAKSPVKEEAKSPAEAKSPVKE SPAEVKSPEKAKSPVKEEAKSPEKAKSPVKE EKEEAKSPEKAKSPVKAEAKSPVKEEAKSPVK KSPEKAKSPVKEEAKSPVKEEAKSPVKEEAKSP KSPVKEEAKTPEKAKSPVKEEAKSPEKAKSP KTLDVKSPEAKTPAKEEARSPADKFPEKAKSEVK EEVKSPEKAKSPLKEDAKAPEKEIPKKEEVK KEEEKPQEVKVKEPPKKAEEEKAPATPKTEI DSKKEEAPKKEAPKPVEEKEAPAKVEVKEDA EKTEVAKKEPDDAKAKEPSKPAEKKEAAPE
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EKEEAKSPEKAKSPVKAEAKSPEKAKSPVK KSPEKAKSPVKEEAKSPKAKSPVKEEAKSI KSPVKEEAKTPEKAKSPVKEEAKSPEKAKSI KTLDVKSPEAKTPAKEEARSPADKFPEKAKSI KTLDVKSPEKAKSPLKEDAKAPEKEIPKKEEVK EEVKSPEKAKSPLKEDAKAPEKEIPKKEEVK KEEEKPQEVKVKEPPKKAEEEKAPATPKTEI DSKKEEAPKKEAPKPKVEEKKEPAVEKPKE EAKKEEAEDKKKVPTPEKEAPAKVEVKEDA
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KSPVKEEAKTPEKAKSPVKEEAKSPEKAKSI KTLDVKSPEAKTPAKEEARSPADKFPEKAKSI EEVKSPEKAKSPLKEDAKAPEKEIPKKEEVK KEEEKPQEVKVKEPPKKAEEEKAPATPKTEI DSKKEEAPKKEAPKPKVEEKKEPAVEKPKE EAKKEEAEDKKKVPTPEKEAPAKVEVKEDA
KTLDVKSPEAKTPAKEEARSPADKFPEKAK EEVKSPEKAKSPLKEDAKAPEKEIPKKEEVK KEEEKPQEVKVKEPPKKAEEEKAPATPKTE DSKKEEAPKKEAPKPKVEEKKEPAVEKPKE EAKKEEAEDKKKVPTPEKEAPAKVEVKEDA EKTEVAKKEPDDAKAKEPSKPAEKKEAAPE
EEVKSPEKAKSPLKEDAKAPEKEIPKKEEVK KEEEKPQEVKVKEPPKKAEEEKAPATPKTEI DSKKEEAPKKEAPKPKVEEKKEPAVEKPKE EAKKEEAEDKKKVPTPEKEAPAKVEVKEDA EKTEVAKKEPDDAKAKEPSKPAEKKEAAPE
KEEEKPQEVKVKEPPKKAEEEKAPATPKTEJ DSKKEEAPKKEAPKVEEKKEPAVEKPKE EAKKEEAEDKKKVPTPEKEAPAKVEVKEDA EKTEVAKKEPDDAKAKEPSKPAEKKEAAPE
DSKKEEAPKKEAPKPKVEEKKEPAVEKPKE EAKKEEAEDKKKVPTPEKEAPAKVEVKEDA EKTEVAKKEPDDAKAKEPSKPAEKKEAAPE
EAKKEEAEDKKKVPTPEKEAPAKVEVKEDA EKTEVAKKEPDDAKAKEPSKPAEKKEAAPE
EKTEVAKKEPDDAKAKEPSKPAEKKEAAPE
TKEEKAKKPEEKPKTEAKAKEDDKTLSKEP
KAEKAEKSSSTDQKDSKPPEKATEDKAAKO
3456 A 258 1463 YLSFIPGHASKSAPMNGHCFAENGPSQKSSL
IPPSENLGPHEEDQVVCGFKKLTVNGVCAST TPIKNSPSLFPCAPLCERGSRPLPPLPISEALSI
DCEVEFLTSSDTDFLLEDSTLSDFKYDVPG*
RGCGQINYAYFDTPAVSAADLSYVSDQNG\\
DPNPPPPQTHRRLRRSHSGPAGSFNKPAIRIS
HRASPNSDEDKPEVPPRVKPDYRR
EVTSSTYSDEDRPPKVPPREPLSPSNSRTPSPI
SYLNGVMPPTQSFAPDPKYVSSKALQRQNS
ASKVPCILPIIENGKKVSSTHYYLLPERPPYL
EKFFREAKKKNGGAQIQPLPADCGISSATEK
KTKMDLGGHVKRKHLSYVGTP
3457 A 2 4869 FILSSSSSASSEHFHHHYSFGNWWPGSFKGH
LPFYQRCHQHYDLSYRNKDVRSTVSHYQRE
SAVYTQGSTAYSSRSSAAHRRESEAFRRASA
QQQASQHALSSEVSRKAASAYDYGSSHGLT
LLLDDYSSKLSPKPKRAKHSLLSGEEKENLP
MVPIFSGRQKHVSGITDTEEERIKEAAAYIAG
LASEEGITTPKQSTASKQTTASKQSTASKQS
QSTASRQSTASRQSVVSKQATSALQQEETSE
RKVVIRGKAERLSLRKTLEETETYHAKLNEI
HAPEFIIKPRSHTVWEKENVKLHCSIAGWPE
HAPEFIIKPRSHTVWEKENVKLHCSIAGWPE
HAPEFIIKPRSHTVWEKENVKLHCSIAGWPE TWYKNQVPINVHANPGKYIIESRYGMHTLE

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			I No. 19 4 A S	I A S TO S TO S TO S TO S TO S TO S TO S
SEQ ID	Method	Predicted	Predicted end nucleotide	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine,
NO:		beginning nucleotide	location	I=Isoleucine, K=Lysine, L=Leucine, M=Methionine,
		location	corresponding	N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine,
		corresponding	to last amino	T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine,
		to first amino	acid residue of	X=Unknown, *=Stop codon, /=possible nucleotide deletion,
		acid residue of peptide	peptide sequence	\─possible nucleotide insertion
İ		sequence	sequence	
	 			WYRNGVPLSPSKWVQTLWSGERATLTFSHLNKE
				DEGLYTIRVRMGEYYEQYSAYVFVRDADAEIEG
				APAAPLDVKCLEANKDYIIISWKQPAVDGGSPIL
				GYFIDKCEVGTDSWSQCNDTPVKFARFPVTGLIE
				GRSYIFRVRAVNKMGIGFPSRVSEPVAALDPAEK
	l			ARLKS/PPLSTLDWT\VIVTEEEPSEGIVPGPPTDLS
	İ	1		VTEATRSYVVLSWKPPGQRGHEGIMYFVEKCEA
}				GTENWQRVNTELPVKSPRFALFDLAEGKSYCFR
				VRCSNSAGVGEPSEATEVTVVGDKLDIPKAPGKI
				IPSRNTDTSVVVSWEESKDAKELVGYYIEANVA
				GSGKWEPCNNPVKTHRFTCHGLVTGQSYIFRV
		j		RAVNAAGLSEYSQDSEAIEVKAAIAPPSPPCDITC
				LESFRDSMVLGWKQPDKIGGAEITGYYVNYREV
				IDGVPGKWREANVKAVSEEAYKISNLKENMVY
	1			OFOVAAMNMAGLGAPSAVSECFKCEEWTIAVP
				GPPHSLKCSEVRKDSLVLQWKPPVHSGRTPVTG
1		į		YFVDLKEAKAKEDQWRGLNEAAIKNVYLKVRG
				LKEGVSYVFRVRAINQAGVGKPSDLAGPVVAET
				RPGTKEVVVNVDDDGVISLNFECDKMTPKSEFS
			1	WSKDYVSTEDSPRLEVESKGNKTKMTFKDLGM
		1		
				DDLGIYSCDVTDTDGIASSYLIDEELKRLLALSH
				EHKFPTVPVKSELAVEILEKGQVRF\WMQAEKLS
				GNAKVNYIFNEKGIFEGPKYKMHIDRNTGIIEMF
ļ	1	1	1	MEKLQDEDEGTYTFQLQDGKATNHSTVVLVGD
			İ	VFKKLQKEAEFQRQEWIRKQGPHFVEYLSWEVT
		¥.		GECNVLLKCKVANIKKETHIVWYKDEREISVDE
	ļ			KHDFKDGICTLLITEFSKKDAGIYEVILKDDRGK
				DKSRLKLVDEAFKELMMEVCKKIALSATDLKIQ
				STAEGIQLYSFVTYYVEDLKVNWSHNGSAIRYSD
				RVKTGVTGEQIWLQINEPTPNDKGKYVMELFDG
				KTGHQKTVDLSGQAYDEAYAEFQRLKQAAIAEK
		,	1	NRARVLGGLPDVVTIQEGKALNLTCNVWGDPPP
	1			EVSWLKNEKALASDDHCNLKFEAGRTAYFTING
				VSTADSGKYGLVVKNKYGSETSDFTVSVFIPEEE
				ARMAALESLKGGKKAK
3458	A .	3963	827	LSRSSSDNNTNTLGRNVMSTATSPLMGAQSFPNL
				TTPGTTSTVTMSTSSVTSSSNVATATTVLSVGQS
				LSNTLTTSLTSTSSESDTGQEAEYSLYDFLDSCRA
				STLLAELDDDEDLPEPDEEDDENEDDNQEDQEY
)		EEVMILRRPSLQRRAGSRSDVTHHAVTSQLPQVP
]"		AGAGSRPIGEQEEEEYETKGGRRRTWDDDYVLK
				RQFSALVPAFDPRPGRTNVQQTTDLEIPPPGTPHS
				ELLEEVECTPSPRLALTLKVTGLGTTREVELPLTN
				FRSTIFYYVQKLLQLSCNGNVKSDKLRRIWEPTY
		}	1	TIMYREMKDSDKEKENGKMGCWSIEHVEQYLG
		1		TDELPKNDLITYLQKNADAAFLRHWKLTGTNKS
			1	IRKNRNCSQLIAAYWDLG\EHGTK\SGLNQGAIST
		1	1	LQSSDILNLTKEQPQAKAGNGQNSCGVEDVLQL
		1		LRILYIVASDPYSRISQEDGDEQPQFTFPPDEFTS/
				KKITTKILQQIEEPLALASGALPDWCEQLTSKCPF
		[LIPFETRQLYFTCTAFGASRAIVWLQNRREATVE
			ĺ	RTRTTSSVRRDDPGEFRVGRLKHERVKVPRGESL
		İ	1	MEWAENVMQIHADRKSVLEVEFLGEEGTGLGPT
				LEFYALVAAEFQRTDLGAWLCDDNFPDDESRHV
				DLGGGLKPPGYYVQRSCGLFTAPFPQDSDELERI
L	ــــــــــــــــــــــــــــــــــــــ			7

SEQ ID NO:	Method	Predicted beginning nucleotide location corresponding to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, I=Icucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion
				TKLFHFLGIFLAKCIQDNRLVDLPISKPFFKLMCM GDIKSNMSKLIYESRGDRDLHCTESQSEASTEEG HDSLSVGSFEEDSKSEFILDPPKPKPPAWFNGILT WEDFELVNPHRARFLKEIKDLAIKRRQILSNKGL SEDEKNTKLQELVLKNPSGSGPPLSIEDLGLNFQF CPSSRIYGFTAVDLKPSGEDEMITMDNAEEYVDL MFDFCMHTGIQKQMEAFRDGFNKVFPMEKLSSF SHEEVQMILCGNQSPSWAAEDIINYTEPKLGYTR DSPGFLRFVRVLCGMSSDERKAFLQFTTGCSTLP PGGLANLHPRLTVVRKVDATDASYPSVNTCVHY LKLPEYSSEEIMRERLLAATMEKGFHLN
3459	A	88	603	SCGPRGLASLGLGFSGRCDDQNKGRS\DGPEAQA EACSGERTYQELLVNQNPIAQPLASRRLTRKLYK CIKKAVKQKQIRRGVKEVQKFVNKGEKGIMVLA GDTLPIEVYCHLPVMCEDRNLPYVYIPSKTDLGA AAGSKRPTCVIMVKPHEEYQEAYDECLEEVQSL PLPL
3460	Α	139	1997	QVTNMSDKSELKAELERKKQRLAQIREEKKRKE EERKKKETDQKKEAVAPVQEESDLEKKRREAEA LLQSMGLTPESPIVPPPMSPSSKSVSTPSEAGSQD SGDGAVGSRRGPIKLGMAKITQVDFPPREIVTYT KETQTPVMAQPKEDEEEDDDVVAPKPPIEPEEEK TLKKDEEN\DSKAPPHELTEEEKQQILHSEEFLSFF DHSTRIVERALSEQINIFFDYSGRDF/ENDKEGEIQ AGAKLSLNRQFF\DER\WSKASGWVSCLDWSSQ YP\ELLVASYNNNEDAPHEPDGVALVWNMKYK KTTPEYVFHCQSAVMSATFAKFHPNLVVGGTYS GQIVLWDNRSNKRTPVQRTPLSAAAHTHPVYCV NVVGTQNAHNLISISTDGKICSWSLDMLSHPQDS MELVHKQSKAVAVTSMSFPVGDVNNFVVGSEE GSVYTACRHGSKAGISEMFEGHQGPITGIHCHAA VGAVDFSHLYVTSSFDWTVKLWTTKNNKPLYSF EDNAGYVYDVMWSPTHPALFACVDGMGRLDL WNLNNDTEVPTASISVEGNPALNRVRWTHSGRE IAVGDSEGQIVIYDVGEQIAVPRNDEWARFGRTL AEINANRADAEEEAATRIPA
3461	A	139	1997	QVTNMSDKSELKAELERKKQRLAQIREEKKRKE EERKKKETDQKKEAVAPVQEESDLEKKRREAEA LLQSMGLTPESPIVPPPMSPSSKSVSTPSEAGSQD SGDGAVGSRRGPIKLGMAKITQVDFPPREIVTYT KETQTPVMAQPKEDEEEDDDVVAPKPPIEPEEEK TLKKDEEN\DSKAPPHELTEEEKQQILHSEEFLSFF DHSTRIVERALSEQINIFFDYSGRDF/ENDKEGEIQ AGAKLSLNRQFF\DER\WSKASGWVSCLDWSSQ YP\ELLVASYNNNEDAPHEPDGVALVWNMKYK KTTPEYVFHCQSAVMSATFAKFHPNLVVGGTYS GQIVLWDNRSNKRTPVQRTPLSAAAHTHPVYCV NVVGTQNAHNLISISTDGKICSWSLDMLSHPQDS MELVHKQSKAVAVTSMSFPVGDVNNFVVGSEE GSVYTACRHGSKAGISEMFEGHQGPITGIHCHAA VGAVDFSHLYVTSSFDWTVKLWTTKNNKPLYSF EDNAGYVYDVMWSPTHPALFACVDGMGRLDL WNLNNDTEVPTASISVEGNPALNRVRWTHSGRE LAVGDSEGQIVIYDVGEQIAVPRNDEWARFGRTL AEINANRADAEEEAATRIPA

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SEQ ID NO:	Method	Predicted beginning nucleotide location corresponding to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion
3462			2643	TAPEFSRSTHASAHASVARVLRNREIAQLKKEQR RQEFQIRALESQKRQQEMVLRRKTQEVSALRRL AKPMSERVAGRAGLKPPMLDSGAEVSASTTSSE AESGARSVSSIVRQWNRKINHFLGDHPAPTVNGT RPARKKFQKKGASQSFSKAARLKWQSLERRIDI VMQRMTIVNLEADMERLIKKREELFLLQEALRR KRERLQAESPEEEKGLQELAEEIEVLAANIDYIND GITDCQATIVQLEETKEELDSTDTSVVISSCSLAE ARLLLDNFLKASIDKGLQVAQKEAQIRLLEGRLR QTDMAGSSQNHLLLDALREKAEAHPELQALIYN VQQENGYASTDEEISEFSEGSFSQSFTMKGSTSH DDFKFKSEPKLSAQMKAVSAECLGPPLDISTKNI TKSLASLVEIKEDGVGFSVRDPYYRDRVSRTVSL PTRGSTFPRQSRATETSPLTRRKSYDRGQPIRSTD VGFTPPSSPPTRPRNDRNVFSRLTSNQSQGSALD KSDDSDSSLSEVLRGIISPVGGAKGARTAPLQCV SMAEGHTKPILCLDATDELLFTGSKDRSCKMWN LVTGQEIAALKGHPNNVVSIKYCSHSGLVFSVST SYIKVWDIRDSAKCIRTLTSSGQVISGDACAATST RAITSAQGEHQINQIALSPSGTMLYAASGNAVRI WELSRFQPVGKLTGHIGPVMCLTVTQTASQHDL VVTGSKDHYVKMFELGECVTGTIGPTHNFEPPH YDGIECLAIQGDILFSGSRDNGIKKWDLDQQELIQ QIPNAHKDWVCALAFIPGRPMLLSACRAGVIKV WNVDNFTPIGEIKGHDSPINAICTNAKHIFTASSG CRVKVWNYVPGLTPCLPRRVLAIKGRATTLP
3463	A	198	3146	SGEPRPEPGNMATCIGEKIEDFKVGNLLGKGSFA GVYRAESIHTGLEVAIKMIDKKAMYKAGMVQR VQNEVKIHCQLKHPSILELYNYFEDSNYVYLVLE MCHNGEMNRYLKNRVKPFSENEARHFMHQIITG MLYLHSHGILHRDLTLSNLLLTRNMNIKIADFGL ATQLKMPHEKHYTLCGTPNYISPEIATRSAHGLE SDVWSLGCMFYTLLIGRPPFDTDTVKNTLNKVV LADYEMPTFLSIEAKDLIHQLLRRNPADRLSLSSV LDHPFMSRNSSTKSKDLGTVEDSIDSGHATISTAI TASSSTSISGSLFDKRRLLIGQPLPNKMTVFPKNK SSTDFSSSGDGNSFYTQWGNQETSNSGRGRVIQD AEERPHSRYLRRAYSSDRSGTSNSQSQAKTYTM ERCHSAEMLSVSKRSGGGENEERYSPTDNNANIF NFFKEKTSSSSGSFERPDNNQALSNHLCPGKTPFP FADPTPQTETVQQWFGNLQINAHLRKTTEYDSIS PNRDFQGHPDLQKDTSKNAWTDTKVKKNSDAS DNAHSVKQQNTMKYMTALHSKPEIIQQECVFGS DPLSEQSKTRGMEPPWGYQNRTLRSITSPLVAHR LKPIRQKTKKAVVSILDSEEVCVELVKEYASQEY VKEVLQISSDGNTITIYYPNGG'RGFPLA'DRPPSP TVDNISR'YSF\DNLPEKYWRKYQYASRFVQLVRS KSPKITYFTRYAKCILMENSPGADFEVWFYDGV KIHKTEDFIQVIEKTGKSYTLKSESEVNSLKEEIK MYMDHANEGHRICLALESIISEEERKTRSAPFFPII IGRKPGSTSSPKALSPPPSVDSNYPTRDRASFNRM VMHSAASPTQAPILNPSMVTNEGLGLTTTASGTD ISSNSLKDCLPKSAQLLKSVFVKNVGWATQ'LTS GAVWVQFNDGSQLVVQAGVSSISYTSPNGQ'TTR 'YGENEKLPDYIKQKLQCLSSILLMFSNPTPNFH

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SEQ ID	Method	Predicted	Predicted end	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid,
NO:		beginning	nucleotide	E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine,
		nucleotide location	location corresponding	N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine,
Í	ľ	corresponding	to last amino	T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine,
		to first amino	acid residue of	X=Unknown, *=Stop codon, /=possible nucleotide deletion,
İ		acid residue of peptide	peptide sequence	∖=possible nucleotide insertion
		sequence	sequence	
3464	A	14	348	AVRTVSGTSLGPRSHSRSPGRCHCFSAVTFSSPRL
				AASEAPDPMEEWDVPQMKKEVESLKYQLAFQR
	ļ			EMASKTIPELLKWIEDGIPKDPFLNPDLMKNNPW
	<u> </u>			VEKGKCTIL
3465	A	5537	405	VRKLDRERVGAWWRGAWARHPRQEAGEHAKR RKGHAETPRGRRKGRAGRSAAAVGELRPARRSL
ŀ				ETSRAAAAMAKDSPSPLGASPKKPGCSSPAAAV
1	}			LENORRELEKLRAELEAERAGWRAERRRFAARE
	1	Į		ROLREEAERERROLADRLRSKWEAQRSRELRQL
				OEEMOREREAEIROLLRWKEAEQRQLQQLLHRE
				RDGVVRQARELQRQLAEELVNRGHCSRPGASEV
				SAAOCRCRLQEVLAQLRWQTDGEQAARIRYLQ
}				AALEVERQLFLKYILAHFRGHPALSGSPDPQAVH
				SLEEPLPQTSSGSCHAPKPACQLGSLDSLSAEVG
				VRSRSLGLVSSACSSSPDGLLSTHASSLDCFAPAC
		1		SRSLDSTRSLPKASKSEERPSSPDTSTPGSRRLSPP
	1			PSPLPPPPPPSAHRKLSNPRGGEGSESQPCEVLTPS
l		1	İ	PPGLGHHELIKLNWLLAKALWVLARRCYTLQEE NKQLRRAGCPYQADEKVKRLKVKRAELTGLAR
				RLADRARELQETNLRAVSAPIPGESCAGLELCQV
				FARQRARDLSEQASAPLAKDKQIEELRQECHLLQ
				ARVASGPCSDLHTGRGGPCTQWLNVRDLDRLQ
				RESQREVLRLQRQLMLQQGNGGAWPEAGGQSA
				TCEEVRRQMLALERELDQRRRECQELGAQAAPA
			Í	RRRGEEAETQLQAALLKNAWLAEENGRLQAKT
				DWVRKVEAENSEVRGHLGRACQERDASGLIAEQ
				LLQQAARGQDRQQQLQRDPQKALCDLHPSWKEI
		1		QALQCRPGHPPEQPWETSQMPESQVKGSRRPKF
	j			HARAEDYAVSQPNRDIQEKREASLEESPVALGES
			ļ	ASVPQVSETVPASQPLSKKTSSQSNSSSEGSMWA TVPSSPTLDRDTASEVDDLEPDSVSLALEMGGSA
				APAAPKLKIFMAQYNYNPFEGPNDHPEGELPLTA
		}		GDYIYIFGDMDEDGFYEGELEDGRRGLVPSNFVE
				QIPDSYIPGCLPAKSPDLGPSQLPAGQDEALEEDS
				LLSGKAQGVVDRGLCQMVRVGSKTEVATEILDT
				KTEACQLGLLQSMGKQGLSRPLLGTKGVLRMAP
				MQLHLQNVTATSANITWVYSSHRHPHVVYLDD
				REHALTPAGVSCYTFQGLCPGTHYRARVEVRLP
				RDLLQVYWGTMSSTVTFDTLLAGPPYPPLDVLV
				ERHASPGVLVVSWLPVTIDSAGSSNGVQVTGYA
l		ł		VYADGLKVCEVADATAGSTLLEFSQLQVPLTWQ KVSVRTMSLCGESLDSVPAQIPEDFFMCHRWPET
				PPFSYTCGDPSTYRVTFPVCPQKLSLAPPSAKASP
				HNPGSCGEPQAKFLEAFFEEPPRRQSPVSNLGSE
				GECPSSGAGSQAQELAEAWEGCRKDLLFQKSPQ
1				NHRPPSVSDQTGEKENCYQHMGTSKSPAPGFIHL
				RTECGPRKEPCQEKAALERVLRQKQDAQGFTPP
				QLGASQQYASDFHNVLKEEQEALCLDLWGTERR
				EERREPEPHSRQGQALGVKRGCQLHEPSSALCPA
				PSAKVIKMPRGGPQQLGTGANTPARVFVALSDY
				NPLVMSANLKAAEEELVFQKRQLLRVWGSQDT
				HDFYLSECNRQVGNIPGRLVAEMEVGTEQTDRR
				WRSPAQGHLPSVAHLEDFQGLTIPQGSSLVLQGN
				SKRLPLWTPKIMIAALDYDPGDGQMGGQGKGRL
L	1]	ALRAGDVVMVY\GPMDDQGFYYGELGGHRG\L

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Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, _=nossible nucleotide insertion Predicted Predicted end SEQ ID Method nucleotide NO: beginning nucleotide location corresponding location corresponding to first amino to last amino acid residue of \=possible nucleotide insertion acid residue of peptide sequence peptide sequence **VPANLRIKMSSQGH** MSKPPDLLLRLLRGAPRQRVCTLFIIGFKFTFFVSI 3466 A 1 1111

3466	A.		1111	MIYWHVVGEPKEKGQLYNLPAEIPCPTLTPPTPP SHGPTPGNIFFLETSDRTNPNFLFMCSVESAARTH PESHVLVLMKGLPGGNASLPRHLGISLLSCFPNV QMLPLDLRELFRDTPLADWYAAVQGRWEPYLL PVLSDASRIALMWKFGGIYLDTDFIVLKNLRNLT NVLGTQSRYVLNGAFLAFERRHEFMALCMRDFV DHYNGWIWGHQGPQLLTRVFKKWCSIRSLAESR ACRGVTTLPPEAFYPIPWQDWKKYFEDINPEELP RLLSATYAVHVWNKKSQGTRFEATSRALLAQLH ARYCPTTHE/DHENVLVKGPAGHLPNLLLMGHW
3467	A		2175	MAKVILKQSKQCKNLLTCKVAQVCPVCGCLHC YFWWLSGLESRRPSSPLIDIKPIEFGVLSAKKEPIQ PSVLRRTYNPDDYFRKFEPHLYSLDSNSDDVDSL TDEEILSKYQLGMLHFSTQYDLLHNHLTVRVIEA RDLPPPISHDGSRQDMAHSNPYVKICLLPDQKNS KQTGVKRKTQKPVFEERYTFEIPFLEAQRRTLLL TVVDFDKFSRHCVIGKVSVPLCEVDLVKGGHW WKAHDSQFSAPGLPADQQFFADLFSGLVLNPQL LGRVWFASQPASLPVGSLCIDFPRLDIVLRGEYG NLLEAKQQRLVEGEMLFIPARAANLPVNNKPVM LLSLVFAPTWLGLSFYDSRTTSLLHPARQIQLP\SL QRGEGEAMLS\ALTLFSRSPLEQNIIQPLVLSLLHL CGSVVNMPPGNSQPRGDFLYHSICTWVQDNYAQ PLTRESVAQFFNITPNHLSKLFAQHGTMRFIEYVR WVRMAKARMILQKYHLSIHEVAQRCGFPDSDYF CRVFRRQFGMDYVDILQIHRWDYNTPIEETLEAL NDVVKAGKARYIGASSMHASQFAQALELQKQH GWAQFVSMQDHYNLIYREEEREMLPLCYQEGV AVIPWSPLARGRLTRPWGETTARLVSDEVGKNL YKESDENDAQIAERLTGVSEELGATRAQVALAW LLSKPGIAAPIIGTSREEQLDELLNAVDITLKPEQI AELETPYKPHPVVGFK
3468	A	147	3209	ALPLPLPTLYPGMSRRKQRKPQQLISDCEGPSASE NGDASEEDHPQVCAKCCAQFTDPTEFLAHQNAC STDPPVMVIIGGQENPNNSSASSEPRPEGHNNPQ VMDTEHSNPPDSGSSVPTDPTWGPERRGEESSGH FLVAATGTAAGGGGGLILASPKLGATPLPPESTP APPPPPPPPPPPPGVGSGHLNIPLILEELRVLQQRQI HQMQMTEQICRQVLLLGSLGQTVGAPASPSELP GTGTASSTKPLLPLFSPIKPVQTSKTLASSSSSSS SSGAETPKQAFFHLYHPLGSQHPFSAGGVGRSHK PTPAPSPALPGSTDQLIASPHLAFPSTTGLLAAQC LGAARGLEATASPGLLKPKNGSGELSYGEVMGP LEKPGGRHKCRFCAKVFGSDSALQIHLRSHTGER PYKCNVCGNRFTTRGNLKVHFHRHREKYPHVQ MNPHPVPEHLDYVITSSGLPYGMSVPPEKAEEEA ATPGGGVERKPLVASTTALSATESLTLLSTSAGT ATAPGLPAFNKFVLMKAVEPKNKADENTPPGSE GSAISGVAESSTATRMQLSKLVTSLPSWALLTNH FKSTGSFPLPLCARALG\ASPSETSKLQQLVEKID RQGAVAVTSAASGAPTTSAPAPSSSASSGPNQCV ICLRVLSCPRALRLHYGQHGGERPFKCKVCGRAF STRGNLRAHFVGHKASPAARAQNSCPICQKKFT

Predicted end Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, Method Predicted SEO ID E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, NO: beginning nucleotide I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, nucleotide location N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, location corresponding to last amino T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, corresponding to first amino acid residue of X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion acid residue of peptide sequence peptide sequence NAVTLQQHVRMHLGGQIPNGGTALPEGGGAAQ **ENGSEQSTVSGAGSFPQQQSQQPSPEEELSEEEEE EDEEEEDVTDEDSLAGRGSESGGEKAISVRGDS EEASGAEEEVGTVAAAATAGKEMDSNEKTTOOS** SLPPPPPPDSLDQPQPMEQGSSGVLGGKEEGGKP ERSSSPASALTPEGEATSVTLVEELSLQEAMRKEP GESSSRKACEVCGQAFPSQAAL\EEH\QKTHPKEG PLF\TCVFCRQGFLERATLKKHMLLAHHQVQPFA PHGPQNIAALSLVPGCSPSITSTGLSPFPRKDDPTI NLRPLSFALFLGDPNMANLEESFPRGGTRKIHKP 3469 A 3 5664 EKAFQQSVEQDNLFDISTEEGSTKRKKSQKGPAK TKKLKIEKRESSKSAREKFEILSVESLCEGMRILG CVKEVNELELVISLPNGLQGFVQVTEICDAYTKK LNEQVTQEQPLKDLLHLPELFSPGMLVRCVVSSL GITDRGKKSVKLSLNPKNVNRVLSAEALKPGML LTGTVSSLEDHGYLVDIGVDGTRAFLPLLKAQEY IRQKNKGAKLKVGQYLNCIVEKVKGNGGVVSLS VGHSEVSTAIATEQQSWNLNNLLPGLVVKAQVQ KVTPFGLTLNFLTFFTGVVDFMHLDPKKAGTYFS NOAVRACILCVHPRTRVVHLSLRPIFLQPGRPLTR LSCONLGAVLDDVPVQGFFKKAGATFRLKDGVL AYARLSHLSDSKNVFNPEAFKPGNTHKCRIIDYS **QMDELALLSLRTSIEAQYLRYHDIEPGAVVKGT** VLTIKSYGMLVKVGEQMRGLVPPMHLADILMK NPEKKYHIGDEVKCRVLLCDPEAKKLMMTLKKT LIESKLPVITCYADAKPGLQTHGFIIRVKDYGCIV KFYNNVOGLVPKHELSTEYIPDPERVFYTGOVV KVVVLNCEPSKERMLLSFKLSSDPEPKKEPAGHS **OKKGKAINIGQLVDVKVLEKTKDGLEVAVLPHN** IRAFLPTSHLSDHVANGPLLHHWLQAGDILHRVL CLSQSEGRVLLCRKPALVSTVEGGQDPKNFSEIH **PGMLLIGFVKSIKDYGVFIQLPSGLSGLAPKAIMS** DKFVTSTSDHFVEGQTVAAKVTNVDEEKQRMLL SLRLSDCGLGDLAITSLLLLNQCLEELQGVRSLM SNRDSVLIQTLAEMTPGMFLDLVVQEVLEDGSV VFSGGPVPDLVLKASRYHRAGQEVESGQKKKVV ILNVDLLKLEVHVSLHQ\DLV\NRKARKLRKGSE HOAIVOHLEKSFAIASLVETGHLAAFSLTSHLND TFRFDSEKLQVGQGVSLTLKTTEPGVTGLLLAVE **GPAAKRTMRPTQKDSETVDEDEEVDPALTVGTI** KKHTLSIGDMVTGTVKSIKPTHVVVTLEDGIIGCI HASHILDDVPEGTSPTTKLKVGKTVTARVIGGRD MKTFKYLPISHPRFVRTIPELSVRPSELEDGHTAL NTHSVSPMEKIKQYQAGQTVTCFLKKYNVVKK WLEVEIAPDIRGRIPLLLTSLSFKVLKHPDKKFRV GQALRATVVGPDSSKTFLCLSLTGPHKLEEGEVA MGRVVKVTPNEGLTVSFPFGKIGTVSIFHMSDSY SETPLEDFVPQKVVRCYILSTADNVLTLSLRSSRT NPETKSKVEDPEINSIQDIKEGQLLRGYVGSIQPH GVFFRLGPSVVGLARYSHVSQHSPSKKALYNKH LPEGKLLTARVLRLNHQKNLVELSFLPGDTGKPD VLSASLEGQLTKQEERKTEAEERDQKGEKKNQK RNEKKNOKGOEEVEMPSKEKOOPOKPOAOKRG GRECRESGSEQERVSKKPKKAGLSEEDDSLVDV

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SEQ ID NO:	Method	Predicted beginning nucleotide location corresponding to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion
				YYREGKEEAEETNVLPKEKQTKPAEAPRLQLSSG FAWNVGLDSLTPALPPLAESSDSEEDEKPHQATI KKSKKERELEKQKAEKELSRTEEALMDPGRQPE SADDFDRLVLSSPNSSILWLQYMAFHLQATEIEK ARAVAERALKTISFREEQEKLNVWVALLNLENM YGSQESLTKVFERAVQYNEPLKVFLHLADIYAKS EKFQEAGELYNRMLKRFRQEKAVWIKYGAFLLR RSQAAASHRVLQRALECLPSKEHVDVIAKFAQL EFQLGDAERAKAIFENTLSTYPKRTDVWSVYID MTIKHGSQKDVRDIFERVIHLSLAPKRMKFFFKR YLDYEKQHGTEKDVQAVKAKALEYVEAKSSVL ED
3470	A	2334	1226	TAAAPVAPGTMDDATVLRKKGYIVGINLGKGSY AKVKSAYSERLKFNVAVKIIARKKTPTDFVERFL PREMDILATVNHGSIIKTYEIFETSDGRIYIIMELG VQGDLLEFIKCQGALHEDVARKMFRQLSSAVKY CHDLDIVHRDLKCENLLLDKDFNIKLSDFGFSKR CLRDSNGRIILSKTFCGSAAYAAPEVLQSIPYQPK VYDIWSLGVILYIMVCGSMPYDDSDIRKMLRIQK EHRVDFPRSKNLTCECKDLIYRMLQ\PDVS\KRLH IDEILSHSWLQPPKPK\ATSSASFKREGEGKYRAE CKLDTKTGLRPDHRPDHKLGAKTQHRLLVVPEN ENRMEDRLAETSRAKDHHISGAEVGKAST
3471	A	537	148	TERGAPQHPTLPLPSLTPSSVHTGQPKTTPSVILFL PSCEEPQANKATLVCLMNN/FYPGILMVTWKAD GTLITQSVEKTTPSKQSNNKYVASSYLSLTPEQW RSRRSYSCQVMQEGSTVEKSVAPAECS
3472	A	1		DKPTRHKTYLSSSWAKMAAAEGPVGDGELWQT WLPNHVVFLRLREGLKNQSPTEAEKPASSSLPSS PPPQLLTRNVVFGLGGELFLWDGEDSSFLVVRLR GPSGGGEEPALSQYQRLLCINPPLFEIYQVLLSPT QHHVALIGIKGLMVLELPKRWGKNSEFEGGKST VNCSTTPVAERFFTSSTSLTLKHAAWYPSEILDPH VVLLTSDNVIRIYSLREPQTPTNVIILSEAEEESLV LNKGRAYTASLGETAVAFDFGPLAAVPKTLFGQ NGKDEVVAYPLYILYENGETFLTYISLLHSPGN/I WKAVGSIAHAS\AAEDNYGYDACAVLCLPCVPN ILVIATESGMLYHCVVLEGEEEDDHTSEKSWDSR IDLIPSLYVFECVELELALKLASGEDDPFDSDFSC PVKLHRDPKCPSRYHCTHEAGVHSVGLTWIHKL HKFLGSDEEDKDSLQELSTEQKCFVEHILCTKPLP CRQPAPIRGFWIVPDILGPTMICITSTYECLIWPLL STVHPASPPLLCTREDVEVAESPLRVLAETPDSFE KHIRSILQRSVANPAFLKASEKDIAPPPEECLQLLS RATQVFREQYILKQDLAKEEIQRRVKLLCDQKK KQLEDLSYCREERKSLREMAERLADKYEEAKEK QEDIMNRMKKLLHSFHSELPVLSDSERDMKKEL QLIPDQLRHLGNAIKQVTMKKDYQQQKMEKVL SLPKPTIILSAYQRKCIQSILKEEGEHIREMVKQIN DIRNHVNF
3473	A		2272	DKPTRHKTYLSSSWAKMAAAEGPVGDGELWQT WLPNHVVFLRLREGLKNQSPTEAEKPASSSLPSS PPPQLLTRNVVFGLGGELFLWDGEDSSFLVVRLR GPSGGGEEPALSQYQRLLCINPPLFEIYQVLLSPT QHHVALIGIKGLMVLELPKRWGKNSEFEGGKST

SEQ ID Method	Predicted	Predicted end	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid,
NO:	beginning	nucleotide	E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine,
	nucleotide location	location corresponding	I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine,
	corresponding	to last amino	T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine,
	to first amino	acid residue of	X=Unknown, *=Stop codon, /=possible nucleotide deletion, \ -possible nucleotide insertion
	acid residue of peptide	peptide sequence	Phossible indiceonde insertion
	sequence		LANGOTTON A ED PETECOTOL TIL ELLA ANTADOLTI DOLL
			VNCSTTPVAERFFTSSTSLTLKHAAWYPSEILDPH VVLLTSDNVIRIYSLREPQTPTNVIILSEAEEESLV
			LNKGRAYTASLGETAVAFDFGPLAAVPKTLFGQ
			NGKDEVVAYPLYILYENGETFLTYISLLHSPGN/I
			WKAVGSIAHAS\AAEDNYGYDACAVLCLPCVPN
			ILVIATESGMLYHCVVLEGEEEDDHTSEKSWDSR
			IDLIPSLYVFECVELELALKLASGEDDPFDSDFSC
			PVKLHRDPKCPSRYHCTHEAGVHSVGLTWIHKL
			HKFLGSDEEDKDSLQELSTEQKCFVEHILCTKPLP
[CROPAPIRGFWIVPDILGPTMICITSTYECLIWPLL
			STVHPASPPLLCTREDVEVAESPLRVLAETPDSFE
}			KHIRSILQRSVANPAFLKASEKDIAPPPEECLQLLS RATQVFREQYILKQDLAKEEIQRRVKLLCDQKK
			KQLEDLSYCREERKSLREMAERLADKYEEAKEK
1			QEDIMNRMKKLLHSFHSELPVLSDSERDMKKEL
			QLIPDQLRHLGNAIKQVTMKKDYQQQKMEKVL
1. 1			SLPKPTIILSAYQRKCIQSILKEEGEHIREMVKQIN
			DIRNHVNF
3474 A	4344	2550	DRRREPERHVRVKQRTSVLNMLRRLDKIRFRGH
			KRDDFLDLAESPNASDTECSDEIPLKVPRTSPRDS
			EELRDPAGPGTLIMATGVQDFNRTEFDRLNEIKG HLEIALLEKHFLQEELRKLREETNAEMLRQELDR
			ERQRRMELEQKVQEVLKARTEEQMAQQPPKGQ
			AQASNGAERRSQGLSSRLQKWFYERFGEYVEDF
1		1	RFQPEENTVETEEPLSARRLTENMRRLKRGAKPV
			TNFVKNLSALSDWYSVYTSAIAFTVYMNAVWH
			GWAIPLFLAILRLSLNYLIARGWRIQWSIVPEV
			SEPVEPPKEDLTVSEKFQLVLDVAQKAQNLFGK
	1		MADILEKIKNLFMWVQPEITQKLYVALWAAFLA SCFFPYRLVGLAVGLYAGIKFFLIDFIFKRCPRLR
			AKYDTPYIIWRSLPTDPQLKERSSAAVSRRLQTTS
			SRSYVPSAPAGLGKEEDAGRFHSTKKGNFHEIFN
			LTENERPLAVCENGWRCCLINRDRKMPTDYIRN
		ĺ	GVLYVT\ENYLCFESSKSGSSKRNKVIKLVDITDI
			QKYKVLSVLPGSGMGIAVSTPSTQKPLVFGAMV
		1105	HRDEAFETILSQYIKITSAAASGGDS
3475 A	2	1126	TAARRRQKGAAAAAETHGQAKAKSGWLKPYYF IELMESRKDITNQEELWKMKPRRNLEEDDYLHK
			DTGETSMLKRPVLLHLHQTAHADEFDCPSELQH
			TQELFPQWHLPIKIAAIIASLTFLYTLLREVIHPLA
			TSHQQYFYKIPILVINKVLPMVSITLLALVYLPGV
			IAAIVQLHNGTKYKKFPHWLDKWMLTRKQFGL
			LSFFFAVLHAIYSLSYPMRRSYRYKLLNWAYQQ
1			VQQNKEDAL\IEHDVWRMEIYVSLGIVGLAILAL
			LAVTSIPSVSDSLTWREFHYIQSKLGIVSLLLGTIH
			ALIFAWNKWIDIKQFVWYTPPTFMIAVFLPIVVLI
3476 A	143	3191	FKSILFLPCLRKKILKIRHGWEDVTKINKTEICSQL AKAPPTGESSEPEAKVLHTKRLYRAVVEAVHRL
34/0 A	143	3171	DLILCNKTAYQEVFKPENISLRNKLRELCVKLMF
			LHPVDYGRKAEELLWRKVYYEVIQLIKTNKKHI
			HSRSTLECAYRTHLVAGIGFYQHLLLYIQSHYQL
			ELQCCIDWTHVTDPLIGCKKPVSASGKEMDWAQ
			MACHRCLVYLGDLSRYQNELAGVDTELLAERFY
			YQALSVAPQIGMPFNQLGTLAGSKYYNVEAMY
<u> </u>			CYLRCIQSEVSFEGAYGNLKRLYDKAAKMYHQL

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Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, SEQ ID Method Predicted Predicted end nucleotide E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, NO: beginning I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, nucleotide location N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, corresponding location T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, corresponding to last amino X=Unknown, *=Stop codon, /=possible nucleotide deletion, acid residue of to first amino \=possible nucleotide insertion acid residue of peptide sequence peptide sequence KKCETRKLSPGKKRCKDIKRLLVNFMYLQSLLQ PKSSSVDSELTSLCQSVLEDFNLCLFYLPSSPNLS LASEDEEEYESGYAFLPDLLIFQMVIICLMCVHSL ERAGSKOYSAAIAFTLALFSHLVNHVNIRLQAEL EEGENPVPAFQSDGTDEPESKEPVEKEEEPDPEPP PVTPQVGEGRKSRKFSRLSCLRRRRHPPKVGDDS DLSEGFESDSSHDSARASEGSDSGSDKSLEGGGT **AFDAETDSEMNSQESRSDLEDMEEEEGTRSPTLE** PPRGRSEAPDSLNGPLGPSEASIASNLQAMSTQM **FQTKRCFRLAPTFSNLLLQPTTNPHTSASHRPCV** NGDVDKPSEPASEEGSESEGSESSGRSCRNERSIQ **EKLQVLMAEGLLPAVKVFLDWLRTNPDLIIVCA** OSSQSLWNRLSVLLNLLPAAGELQESGLALCPEV ODLLEGCELPDLPSSLLLPEDMALRNLPPLRAAH RRFNFDTDRPLLSTLEESVVRICCIRSFGHFIARLQ GSILQFNPEVGIFVSIAQSEQESLLQQAQAQFRMA **QEEARRNRLMRDMAQLRLQLEVSQLEGSLQQPK AQSAMSPYLVPDTQALCHHLPVIRQLATSGRFIVI IPRTVIDGLDLLKKEHPGARDGIRYLEAEFKKGN** RYIRCQKEVGKSFERHKLKRQDADAWTLYKILD SCKQLT\LAQGAGEEDPSGMVTIITGLPLDNPSVL SGPMQAALQAAAHASVDIKNVLDFYKQWKEIG MTEPRERRGYSVPPRPEVGTQATEWRVEESNFN 3902 3477 Α KIFLKKDAELGRSNHLPTWDKPEDASWLPQSCL **GGDAVATTGEIHEEKAWKTRALEVGQPAQRDIR** RGELWGKEHGADQAIQETLEDLSSLERTLVVSES SPLGGDCQEVTTLTVKYQVSEEVPSGTVIGKLSQ ELGREERRROAGAAFQVLQLPQALPIQVDSEEGL LSTGRRLDREQLCRQWDPCLVSFDVLATGDLALI HVEIQVLDINDHQPRFPKGEQELEISESASLRTRIP LDRALDPDTGPNTLHTYTLSPSEHFALDVIVGPD **ETKHAELIVVKELDREIHSFFDLVLTAYDNGNPP** KSGTSLVKVNVLDSNDNSPAFAESSLALEIQEDA APGTLLIKLTATDPDQGPNGEVEFFLSKHMPPE\V LDTFSIDAKTGQVILRRPLDYEKNPAYEVDVQAR DLGPNPIPAHCKVLIKVLDVNDNIPSIHVTWASQP SLVSEALPKDSFIALVMADDLDSGNNGLVHCWL SOELGHFRLKRTNGNTYMLLTNATLDREQWPK YTLTLLAQDQGLQPLSAKKQLSIQISDINDNAPVF EKSRYEVSTRENNLPSLHLITIKAHDADLGINGK VSYRIQDSPVAHLVAIDSNTGEVTAQRSLNYEEM AGFEFQVIAEDSGQPMLASSVSVWVSLLDANDN APEVVOPVLSDGKASLSVLVNASTGHLLVPIETP NGLGPAGTDTPPLATHSSRPFLLTTIVARDADSG ANGEPLYSIRSGNEAHLFILNPHTGQLFVNVTNA SSLIGSEWELEIVVEDQGSPPLQTRALLRVMFVTS **VDHLRDSARKPGALSMSMLTVICLAVLLGIFGLI** LALFMSICRTEKKDNRAYNCREAESTYRQQPKR POKHIQKADIHLVPVLRGQAGEPCEVGQSHKDV DKEAMMEAGWDPCLQAPFHLTPTLYRTLRNQG NOGAPAESREVLODTVNLLFNHPRQRNASRENL NLPEPQPATGQPRSRPLKVAGSPTGRLAGDQGSE EAPQRPPASSATLRRQRHLNGKVSPEKESGPRQI LRSLVRLSVAAFAERNPVEELTVDSPPVQQISQLL SLLHQGQFQPKPNHRGNKYLAKPGGSRSAIPDTD

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SEQ ID NO:	Method	Predicted beginning nucleotide location corresponding to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion
		- Sequence		GPSARAGGQTDPEQEEGPLDPEEDLSVKQLLEEE
	*** -			LSSLLDPSTGLALDRLSAPDPAWMARLSLPLTTN YRDNVISPDAAATEEPRTFQTFGKAEAPELSPTG TRLASTFVSEMSSLLEMLLEQRSSMPVEAASEAL
				RRLSVCGRTLSLDLATSAASGMKVQGDPGGKTG
3478	A	13	1620	TEGKSRGSSSSSRCL TLPPPGNSGCHRLCFPEFEFLQVTKMEFSGRKWR
34/0	A			KLRLAGDQRNASYPHCLQFYLQPPSENISLIEFEN LAIDRVKLLKSVENLGVSYVKGTEQYQSKLESEL RKLKFSYRENLEDEYEPRRRDHISHFILRLAYCQS EELRRWFIQQEMDLLRFRFSILPKDKIQDFLKDSQ LQFEAISDEEKTLREQEIVASSPSLSGLKLGFESIY KIPFADALDLFRGRKVYLEDGFAYVPLKDIVAIIL NEFRAKLSKALALTARSLPAVQSDERLQPLLNHL SHSYTGQDYSTQGNVGKISLDQIDLLSTKSFPPC MRQLHKALRENHHLRHGGRMQYGLFLKGIGLT LEQALQFWKQEFIKGKMDPDKFDKGYSYNIRHS FGKEGKRTDYTPFSCLKIILSNPPSQGDYHGCPFR HSDPELLKQKLQSYKISPGGISQILDLVKGTHYQ V\ACQKYFEMIHTVDDCGFS\LSHPNQYFCESQRI LNGGKDIKKEPIQPETPQPKPSVQKTKDASSALA SLNSSLEMDMEGLEDYFSEDS
3479	A	698	138	RPELELWRLRSRSWRPLGVPRRCHRRNWKEPVR
				AQPLSVTVWAPRCQRP/QPPAPEPSSPNAAVPEAI PTPRAAASAALELPLGPAPVSVAPQAEAEARSTP GPAGSRLGPETFRQFRQFRYQDAAGPREAFRQL REL/SPRQWLRPDI\RTKEQ\IVEMLVQEQLLAILP EAARARRIRRRTDVRITG
3480	Α	117	2226	RRGSRSRGPFAEPAAPGGLCSSSEEKTEEGGMAV
				GLCKAMSQGLVTFRDVALDFSQEEWEWLKPSQ KDLYRDVMLENYRNLVWLGLSISKPNMISLLEQ GKEPWMVERKMSQGHCADWESWWEIEELSPK WFIDEDEISQEMVMERLASHGLECSSFREAWKY KGEFELHQGNAERHFMQVTAVKEISTGKRDNEF SN/IWEKHTPEISIFNTTES\PTIQQVHKFDIYDKLF PQNSVIIEYKRLHAEKESLIGNECEEFNQSTYLSK DIGIPPGEKPYESHDFSKLLSFHSLFTQHQTTHFG KLPHGYDECGDAFSCYSFFTQPQRIHSGEKPYAC NDCGKAFSHDFFLSEHQRTHIGEKPYECKECNKA FRQSAHLAQHQRIHTGEKPFACNECGKAFSRYAF LVEHQRIHTGEKPYECKECNKAFRQSAHLNQHQ RIHTGEKPYECNQCGKAFSRIALTLHQRIHTGE KPFKCSECGKTFGYRSHLNQHQRIHTGEKPYECI KCGKFFRTDSQLNRHHRIHTGERPFECSKCGKAF SDALVLIHHKRSHAGEKPYECNKCGKAFSCGSY LNQHQRIHTGEKPYECSECGKAFHQILSLRLHQRI HAGEKPYKCNESQRVRRSELAVSRGLTTKPADT GPDSTLNAAKVAEPARAGTEAALRPALSVAESA TSLGPLHQGRRFPEAPAAHPGGTGFTVCAS
3481	A	2	1522	ASRHGMTPGALLMLLGALGPPLAPGVRGSEAEG RLREKLFSGYDSSVRPAREVGDRVRVSVGLILAQ LISLNEKDEEMSTKVYLDLEWTDYRLSWDPAEH DGIDSLRITAESVWLPDVVLLNNNDGNFDVALDI SVVVSSDGSVRWQPPGIYRSSCSIQVTYFPFDWQ NCTMVFSSYSYDSSEVSLQTGLGPDGQGHQEIHI

SEQ ID NO:	Method .	Predicted beginning nucleotide location corresponding to first amino acid residue of peptide	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \;
		sequence		HEGTFIENGQWENIHKPSRLIQPPGDPRGGREGQ RQEVIFYLIIRRKPLFYLVNVIAPCILITLLAIFVFY LPPDAGEKMGLSIFALLTLTVFLLLLADKVPETSL SVPIIIKYLMFTMVLVTFSVILSVVVLNLHHRSPH THQMPLWVRQIFIHKLPLYLRLKRPKPERDLMPE PPHCSSPGSGWGRGTDEYFIRKPPSDFLFPKPNRF QPELSAPDLRRFIDGPNRAVALLPELREVVSSISYI ARQLQEQEDHDALKEDWQFVAMVVDRLFLWTF IIFTSVGTL\VIFLDATYHLPPPDPFP
3482	A	1273	172	ERWDSGGADAEWYALADWTAVWLPRSDFYTR LQTGEGHVPALRLPAGMPPDSPRELVPKQAPCSP SDPALPWTLGHGNQPPAVVPEPQGPMGPAGVAA RPGRFFGVYLLYCLNPRYRVR\VYVGFTVNTARR VQQHNGGRKKGGA\GRTSGRGPWEMVLVVHGF PSSVAALRFEWA WQHPHASRRLAHVGPRLRGET AFAFHLRVLAHMLRAPPWARLPLTLRWVRPDLR QDLCLPPPPHVLLAFGPPPAQVPRPQRRAGPFD DAEPEPDQGDPGACCSLCAQTIQDEEGPLCCPHP GCLLRAHVICLAEEFLQEEPGQLLPLEGQCPCCE KSLLWGDLIWLCQMDTEKEVEDSELEEAHWTD LLET
3483	A	230		WRPWPCIDTSWNLQVAARTLRVSSAQCGLVPT MARVESPVPAARASLTGSCVLGQAMPLRGGAGP SPASHGPTHGPSDPRTCLPGRGAGGMRPHGRGA LGCCGLCSFYTCHGAAGDEIMHQDIVPLCAADIQ DQLKKRFAYLSGGRGQDGSPVITFPDYPAFSEIPD KEFQNVMTYLTSIPSLQDAGIGFILVIDRRDKW TSVKASVLRIAASFPANLQLVLVLRPTGFFQRTLS DIAFKFNRDDFKMKVPVIMLSSVPDLHGYIDKSQ LTEDLGGTLDYCHSRWLCQRTAIESFALMVKQT AQMLQSFGTELAETELPNDVQST\SSVLCAHTEK KDKAKEDLRLALKEGHSVLESLRELQAEGSEPSV NQDQLDNQATVQRLLAQLNETEAAFDEFWAKH QKLEQCLQLRHFEQGFREVKAILDAASQKIATF TDIGNSLAHVEHLLRDLANFQEKSGVFVERARA LSLTASSFIGNKHYAVDSIRPKCQELRHLCDQFSA EIARRRGLLSKSLELHRRLETSMKWCDEGIYLLA SQPVDKCQSQDGAEAALQEIEKFLETGAENKIQE LNAIYKEYESILNQDLMEHVRKVFQKQASMEEV FHRRQASLKKLAARQTRPVQPVAPRPEALAKSP CPSPGIRRGSENSSSEGGALRRGPYRRAKSEMSES RQGRGSAGEEEESLAILRRHVMSELLDTERAYVE ELLCVLEGYAAEMDNPLMAHLLSTGLHNKKDV LFGNMEEIYHFHNRIFLRELENYTDCPELVGRCF LERMEDFQIYEKYCQNKPRSESLWRQCSDCPFFQ ECQRKLDHKLSLDSYLLKPVQRITKYQLLLKEM LKYSRNCEGAEDLQEALSSILGILKAVNDSMHLI AITGYDGNLGDLGKLLMQGSFSVWTDHKRGHT KVKELARFKPMQRHLFLHEKAVLFCKKREENGE GYEKAPSYSYKQSLNMAAVGITENVKGDAKKFE IWYNAREEVYIVQAPTPEIKAAWVNEIRKVLTSQ LQACREASQHRALEQSQSLPLPAPTSTSPSRGNSR NIKKLEERKTDPLSLEGYVSSAPLTKPPEKGKGW SKTSHSLEAPEDDGGWSSAEEQINSSDAEEDGGL GPKKLVPGKYTVVADHEKGGPDALRVRSGDVV

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SEQ ID NO:	Method	Predicted beginning nucleotide location corresponding to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanlne C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion
				ELVQEGDEGLW
3484	A	208	6103	ELVQEGDEGLW VTMAQQAADKYLYVDKNFINNPLAQADWAAK KLVWVPSDKSGFEPASLKEEVGEEAIVELVENGK KVKVNKDDIQKMNPPKFSKVEDMAELTCLNEAS VLHNLKERYYSGLIYTYSGLFCVVINPYKNLPIYS EEIVEMYKGKKRHEMPPHIYAITDTAYRSMMQD REDQSILCTGESGAGKTENTKKVIQYLAYVASSH KSKKDQGELERQLLQANPILEAFGNAKTVKNDN SSRFGKFIRINFDVNGYIVGANIETYLLEKSRAIRQ AKEERTFHIFYYLLSGAGEHLKTDLLLEPYNKYR FLSNGHVTIPGQQDKDMFQETMEAMRIMGIPEEE QMGLLRVISGVLQLGNIVFKKERNTDQASMPDN TAAQKVSHLLGINVTDFTRGILTPRIKVGRDYVQ KAQTKEQADFAIEALAKATYERMFRWLVLRINK ALDKTKRQGASFIGILDIAGFEIFDLNSFEQLCINY TNEKLQQLFNHTMFILEQEEYQREGIEWNFIDFG LDLQPCIDLIEKPAGPPGILALLDEECWFPKATDK SFVEKVMQEQGTHPKFQKPKQLKDKADFCIIHY AGKVDYKADEWLMKNMDPLNDNIATLLHQSSD KFVSELWKDVDRIIGLDQVAGMSETALPGAFKT RKGMFRTVGQLYKEQLAKLMATLRNTNPNFVR CIIPNHEKKAGKLDPHLVLDQLRCNGVLEGIRICR QGFPNRVVFQEFRQRYEILTPNSIPKGFMDGKQA CVLMIKALELDSNLYRIGQSKVFFRAGVLAHLEE ERDLKITDVIIGFQACCRGYLARKAFAKRQQQLT AMKVLQRNCAAYLKLRNWQWWRLFTKVKPLL QVSRQEEEMMAKEEELVKVREKQLAAENRLTE METLQSQLMAEKLQLQEQLQAETELCAEAEELR ARLTAKKQELEEICHDLEARVEEEEERCQHLQA EKKKMQQNIQELEEQLEEESARQKLQLEKVTT EAKLKKLEEEQIILEDQNCKLAKEKKLLEDRIAEF TTNLTEEEKSKSLAKLKNKHEAMITDLEERLRR EEKQRQELEKTRRKLEGDSTDLSDQIAELQAQUA ELKMQLAKKEEELQAALARVEEEAAQKNMALK KIRELESQISELQEDLKCERVASRNKAEKQKRDLG EELALKTELEDTLDSTAAQQELRSKREQEVNIL KKTLEEEAKTHEAQIQEMRQKHSQAVEELAEQL EQTKRVKANLEKAKQTLENERGELANEVKVLLQ GKGDSEHKRKKVEAQLQELQVKFNEGERVRTEL ADKVTKLQVELDNVTGLLSQSDSKSSKLTKDFS ALESQLQDTQELLQEENRQKLSSTKLKQVEDE KNSVFRQLEEEEEEAKHNLEKQIATLHAQVADM KKKMEDSVGCLETAEEVKRKLQKDLEGLSQRHE EKVAAYDKLEKTKTRLQQELDDLLVDLDHQRQ SACNLEKKKKVEAQLQELQVKFNEGERVRTEL ADKVTKLQVELDNVTGLLSQSDSKSSKLTKDFS ALESQLQDTGELLQEENRQKLSSTKKAVEEDRA EEKARREKTKALSLARALEEAMEQKAELERINK GPRTEMEDLMSSKDDVGKSVHELEKSKRAIEQQ VEEMKTQLEEEDELQATEDAKLRLEVNLQAM KAQFERDLQGRDEQSEEKKKQLVRQVREMEAE LEDERKCRSMAVAARKKLEMDLKDLEAHIDSA NKNRDEAIKQLRKLLQAQMKDCMRELDDTRASR EEILAQAKENEKKLKSMEAEMIQLQELAAAER AKRQAQERDELADEIANSSGKGALALEEKRIL
				EARIAQLEEELEEEQGNTELINDRLKKANLQIDQI NTDLNLERSHAQKNENARQQLERQNKELKVKL

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SEQ ID NO:	Method	Predicted beginning nucleotide location corresponding to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion
	-			QEMEGTVKSKYKASITALEAKIAQLEEQLDNETK ERQAACKQVRRTEKKLKDVLLQVDDERRNAEQ YKDQADKASTRLKQLKRQLEEAEEEAQRANASR RKLQRELEDATETADAMNREVSSLKNKLRRGDL PFVVPRRMARKGAGDGSDEEVDGKADGAEAKP AE
3485	A	2	1782	CSTGVSKAPLTYLMSYGFELGWRKGNRAVACR EDRGGESVGMGQESILSQVHWWEAEPVEKTPGR DSEATIMSLRVHTLPTLLGAVVRPGCRELLCLLM ITVTVGPGASGVCPTACICATDIVSCTNKNLSKVP GNLFRLIKRLDLSYNRIGLLDSEWIPVSFAKLNTL ILRHNNITSISTGSFSTTPNLKCLDLSSNKLKTVK NAVFQELKVLEVLLLYNNHISYLDPSAFGGLSQL QKLYLSGNFLTQFPMDLYVGRFKLAELMFLDVS YNRIPSMPMHHINLVPGKQLRGIYLHGNPFVCDV CSLVSLLVFWYRRHFSSVMDFKNDYTCRLWSDS RHSRQVLLLQDSFMNCSDSIINGSFRALGFIHEAQ VGERLMVHCDSKTGNANTDFIWVGPDNRLLEPD KEMENFYVFHNGSLVIESPRFEDAGVYSCIAMNK QRLLNETVDVTINVSNFTVSRSHAHEAFNTAFTT LAACVASIVLVLLYLYLTPCPCKCKTKRQKNML HQSNAHSSILSPGPASDASADERKAGAGKRVVFL EPLKDTAAGQNGKVRLFPSEAVIAEGILKSTRGK SDSDSVNSVFSDTPFVAST
3486	A	357	1173	GDPRETKVFPSRSFARNTVGVSHHQSHLFHTVSR IYVEDKHKILYCEVPKAGCSNWKRILMVLNGLA SSAYNISHNAVHYGKHLKKLDSFDLKGIYTRLDT YTKLVLVRDPMERLVSAFRDKFDHPNSYYHPVF GKAIIKKYRPNACEEALINGSGVKFKEFIHYLLDS HRPVGMDIHWEKVSKLCYPCLINYDFVGKFETL EEDANYFLQMIGAPKELKFPNFKDRHSSDERTNA QVVRQYLKDLTRTERQLIYDFYYLDYLMFNYTT PFL
3487	A		3281	CDKSGAVPFSTTRSPRRPSPRSAGPSLSSVSPRSQ LWASSGLSEEHAAPLLPAWPRHPCPPSLTPGPSM AQGAMRFCSEGDCAISPPRCPRRWLPEGPVPQSP PASMYGSTGSLLRRVAGPGPRGRELGRVTAPCTP LRGPPSPRVAPSPWAPSSPTGQPPPGAQSSVVIFR FVEKASVRPLNGLPAPGGLSRSWDLGGVSPPRPT PALGPGSNRKLRLEASTSDPLPARGGSALPGSRN LVHGPPAPPQVGADGLYSSLPNGLGDPPERLATL FGGPADTGFLNQGDTWSSPREVSSHAQRIARAK WEFFYGSLDPPSSGAKPPEQAPPSPPGVGSRQGS GVAVGRAAKYSETDLDTVPLRCYRETDIDEVLA EREEADSAIESQPSSEGPPGTAYPPAPRPGPLPGP HPSLGSGNEDEDDDEAGGEEDVDDEVFEASEGA RPGSRMPLKSPVPFLPGTSPSADGPDSFSCVFEAI LESHRAKGTSYTSLASLEALASPGPTQSPFFTFEL PPQPPAPRPDPPAPAPLAPLEPDSGTSSAADGPWT QRGEEEEAEARAKLAPGREPPSPCHSEDSLGLGA APLGSEPPLSQLVSDSDSELDSTERLALGSTDTLS NGQKADLEAAQRLAKRLYRLDGFRKADVARHL GKNNDFSKLVAGEYLKFFVFTGMTLDQALRVFL KELALMGETQERERVLAHFSQRYFQCNPEALSSE DGAHTLTCALMLLNTDLHGHNIGKRMTCGDFIG

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SEQ ID NO:	Method	Predicted beginning nucleotide location corresponding to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion
		sequence		NLEGLNDGGDFPRELLKALYSSIKNEKLQWAIDE
	<u> </u>			EELRRFLSELADPNPKVIKRISGGSGSGSSPFLDLT PEPGAAVYKHGALVRKVHADPDCRKTPRGKRG WKSFHGILKGMILYLQKEEYKPGKALSETELKN AISIHHALATRAS\NYSKRPHVFYLRTADWRVFL FQAPSLEQMQSWITRINVVAAMFSAPPFPAAVSS
				QKKFSRPLLPSAATRLSQEEQVRTHEAKLKAMA SELREHRAAQLGKKGRGKEAEEQRQKEAYLEFE KSRYSTYAALLRVKLKAGSEELDAVEAALAQAG STEDGLPPSHSSPSLQPKPSSQPRAQRHSSEPRPG AGSGRRKP
3488	A	441	1968	GTETPHCWGRGTAGLRRELDREERDGPGTATMS FPHFGHPYRGAFQFL\ASASSSTTCCESTLRSVSY VASGSTPAPALCCAP\YDSRLLGSARPELGAALGI YGAPYAAAAAAQSYPGYLPYSPEPPSLYGALNP QYEFKEAAGSFTSSLAQPGAYYPYERTLGQYQY ERYGAVELSGAGRRKNATRETTSTLKAWLNEHR KNPYPTKGEKIMLAIITKMTLTQVSTWFANARR LKKENKMTWAPKNKGGEERKAEGGEEDSLGCL
				TADTKEVTASQEARGLRLSDLEDLEEEEEEEEA EDEEVVATAGDRLTEFRKGAQSLPGPCAAAREG RLERRECGLAAPRFSFNDPSGSEEADFLSAETGSP RLTMHYPCLEKPRIWSLAHTATASAVEGAPPARP RPRSPECRMIPGQPPASARRLSVPRDSACDESSCI PKAFGNPKFALQGLPLNCAPCPRRSEPVVQCQYP SGAEGSGPPAALGVSMQKTPTYRPARQLHTLCH SSLP
3489	A	718	2073	IAAYHKALSYRGHVHANNRGTNNVHFTPPSPS RGILPMNPRNMMNHSQVGQGIGIPSRTNSMSSSG LGSPNRSSPSIICMPKQQPSRQPFTVNSMSGFGMN RNQAFGMNNSLSSNIFNGTDGSENVTGLDLSDFP ALADRNRREGSGNPTPLINPLAGRAPYVGMVTK PANEQSQDFSIHNEDFPALPGSSYKDPTSSNDDSK SNLNTSGKTTSSTDGPKFPGDKSSTTQNNNQQKK GIQVLPDGRVTNIPQGMVTDQFGMIGLLTFIRAA ETDPGMVHLALGSDLTTLGLNLNSPENLYPKFAS PWASSPCRPQDIDFHVPSEYLTNIHIRDKLFFFFS W/TAIKLGRYGEDLLFYLYYMNGGDVLQLLAAV ELFNRDWRYHKEERVWITRAPGMEPTMKTNTY ERGTYYFFDCLNWRKVAKEFHLEYDKLEERPHL PSTFNYNPAQQAF FVAKMATSQYFDFAQGGGPQYSTQAPTLPLPTV
3490	A	2	2833	GASYTGQPTPGMDPAVNPAFPPAAPAGYGGYQP HSGQDFAYGSRPQEPVPTATTMATYQDSYSYGQ SAAARSYEDRPYFQSAALQSGRMTAADSGQPGT QEACGQPSPHGSHSHAQPPQQAPIVESGQPASTL SSGYTYPTATGVQPESSASIVTSYPPPSYNPTCTA YTAPSYPNYDASVYSAASPFYPPAQPPPPPGPPQ QLPPPPAPAGSGSSPRADSKPPLPSKLPRPKAGPR QLQLHYCDICKISCAGPQTYREHLGGQKHRKKE AAQKTGVQPNGSPRGVQAQLHCDLCAVSCTGA DAYAAHIRGSKHQKVFKLHAKLGKPIPTLEPALA TESPPGAEAKPTSPTGPSVCASSRPALAKRPVASK ALCEGPPEPQAAGCRPQWGKPAQPKLEGPGAPT QGGSKEAPAGCSDAQPVGPEYVEEVFSDEGRVL

SEO ID	Method	Predicted	Predicted end	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid,
NO:	Method	beginning	nucleotide	E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine,
	į	nucleotide	location	I=Isoleucine, K=Lysine, L=Leucine, M=Methionine,
		location	corresponding to last amino	N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine,
		corresponding to first amino	acid residue of	X=Unknown, *=Stop codon, /=possible nucleotide deletion,
		acid residue of	peptide	possible nucleotide insertion
		peptide sequence	sequence	
				RFHCKLCECSFNDLNAKDLHVRGRRHRLQYRKK
		-		VNPDLPIATEPSSRARKVLEERMRKQRHLAEERL
				EQLRRWHAERRRLEEEPPQDVPPHAPPDWAQPL
	1		·	LMGRPESPASAPLQPGRRPASSDDRHVMCKHATI
	!			YPTEQELLAVQRAVSHAERALKLVSDTLAEEDR
				GRREEEGDKRSSVAPQTRVLKGVMRVGILAKGL
			<u> </u>	LLRGDRNVRLALLCSEKPTHSLLRRIAQQLPRQL
			1	QMVTEDEYEVSSDPEANIVISSCEEPRMQVTISVT
				SPLMREDPSTDPGVEEPQADAGDVLSPKKCLESL
				AALRHARWFQARASGLQPCVIVIRVLRDLCRRV
				PT\WGALPAWAMELLVEKAVSSAAGPLGPGDAV
				RRVLECVATGTLLTDGPGLQDPCERDQTDALEP
ĺ				MTLQEREDVTASAQHALRMLAFRQTHKVLGMD
				LLPPRHRLGARFRKRQRGPGEGEEGAGEKKRGR
				RGGEGLV
3491	A	2	1321	FVGDGALSGCRRGRAPRVPSMAGSLPPCVVDCG
				TGYTKLGYAGNTEPQFIIPSCIAIRESAKVVDQAQ
1				RRVLRGVDDLDFFIGDEAIDKPTYATKWPIRHGII
				EDWDLMERFMEQVVFKYLRAEPEDHYFLMTEP
				PLNTPENREYLAEIMFESFNVPGLYIAVQAVLAL
			ļ	AASWTSRQVGERTLTGIVIDSGDGVTHVIPVAEG
			ł	YVIGSCIKHIPIAGRDITYFIQQLLREREVGIPPEQS
1				LETAKAIKEKYCYICPDIVKEFAKYDVDPRKWIK
				QYTGINAINQKKFVIDVGYERFLGPEIFFHPEFAN
			ľ	PDFMESISDVVDEVIQNCPIDVRRPLYKNVVLSG
				GSTMFRDFGRRLQRDLKRVVDARLRLSEELSGG\
		Ì	1	RIKPKPVEVQVVTHHMQRYAV\WFGG\SMLASTP
2400	ļ		0004	EFFQVCHTKKDYEEYGPSICRHNPVFGVMS
3492	A	3	2024	PNGVALLHLPGAAVIPNTNYMFQDALGGRSRGS
				REESPAPSRAPASASLWRRLVVVEAKMAAHAAA AAQAAAAQAAHAEAADSWYLALLGFAEHFRTS
1				SPPKIRLCVHCLQAVFPFKPPQRIEARTHLQLGSV
ļ		}	Ì	LYHHTKNSEQARSHLEKAWLISQQIPQFEDVKFE
				AASLLSELYCQENSVDAAKPLLRKAIQISQQTPY
		}		WHCRLLFQLAQLHTLEKDLVSACDLLGVGAEY
			1	ARVVGSEYTRALFLLSKGMLLLMERKLQEVHPL
	}			LTLCGQIVENWQGNPIQKESLRVFFLVLQVTHYL
				DAGOVKSVKPCLKQLQQCIQTISTLHDDEILPSNP
				ADLFHWLPKEHMCVLVYLVTVMHSMQAGYLE
				KAQKYTDKALMQLEKLKMLDCSPILSSFQVILLE
				HIIMCRLVTGHKATALQEISQVCQLCQQSPRLFS
			i	NHAAQLHTLLGLYCVSVNCMDNAEAQFTTALR
				LTNHQELWAFIVTNLASVYIREGNRHQEVV\LYS
				LLERINPDHSFPVSSHCLRAAAFYVRGLFSFFQGR
				YNEAKRFLRETLKMSNAEDLNRLTACSLVLLGHI
				FYVLGNHRESNNMVVPAMQLASKIPDMSVQLW
			1	SSALLRDLNKACGNAMDAHEAAQMHQNFSQQL
			1	LODHIEACSLPEHNLITWTDGPPPVQFQAQNGPN
				TSLASLL
3493	A	3	2024	PNGVALLHLPGAAVIPNTNYMFQDALGGRSRGS
.,,	**		2027	REESPAPSRAPASASLWRRLVVVEAKMAAHAAA
			[AAQAAAAQAAHAEAADSWYLALLGFAEHFRTS
			J	SPPKIRLCVHCLQAVFPFKPPQRIEARTHLQLGSV
ĺ				LYHHTKNSEQARSHLEKAWLISQQIPQFEDVKFE
ĺ				AASLLSELYCQENSVDAAKPLLRKAIQISQQTPY
L	Щ		l	TITANORITI CANTO A DUUTE PRINCIPALITI

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SEQ ID NO:	Method	Predicted beginning nucleotide location corresponding to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \text{\tex{\tex
				WHCRLIFQLAQLHTLEKDLVSACDLLGVGAEY ARVVGSEYTRALFLLSKGMLLLMERKLQEVHPL LTLCGQIVENWQGNPIQKESLRVFFLVLQVTHYL DAGQVKSVKPCLKQLQQCIQTISTLHDDEILPSNP ADLFHWLPKEHMCVLVYLVTVMHSMQAGYLE KAQKYTDKALMQLEKLKMLDCSPILSSFQVILLE HIIMCRLVTGHKATALQEISQVCQLCQQSPRLFS NHAAQLHTLLGLYCVSVNCMDNAEAQFTTALR LTNHQELWAFIVTNLASVYIREGNRHQEVVLYS LLERINPDHSFPVSSHCLRAAAFYVRGLFSFFQGR YNEAKRFLRETLKMSNAEDLNRLTACSLVLLGHI FYVLGNHRESNNMVVPAMQLASKIPDMSVQLW SSALLRDLNKACGNAMDAHEAAQMHQNFSQQL LQDHIEACSLPEHNLITWTDGPPPVQFQAQNGPN TSLASLL
3494	A		1615	VLRGQRGPAGGLAEERRRGRNEWRIHDVTTAPF PGLVQRRSRLLIVSQVRYFLKNKVSPDLCNEDGL TALHQCCIDNFEEIVKLLLSHGANVNAKDNELW TPLHAAATCGHINLVKILVQYGADLLAVNSDGN MPYDLCEDEPTLDVIETCMAYQGITQEKINEMRV APEQQMIADIHCMIAAGQDLDWIDAQGATLLHI AGANGYLRAAELLLDHGVRVDVKDWDGWEPL HAAAFWGQMQMAELLVSHGAN\LNARTSMDE MPIDLCEEEEFKVLLLELK\HKHDVIMKSQLRHK SSLSRRTSHRQAS/SVGKVVRRTQPVGTGPNL\YR KEYE/GEEAILWQRSA\AEDQRTSTYNGDIRET\R TDQENKDPNPRLEK\PVLLSEFPTKIPRGELDMPV ENGLRAPVSAYQYALANGDVWKVHEVPDYSM AYGNPGVADATPPWSSYKEQSPQTLLELKRQRA AAKLLSHPFLSTHLGSSMARTGESSSEGKAPLIG GRTSPYSSNGTSVYYTVTSGDPPLLKFKAPIEEM EEKVHGCCRIS
3495	A	327	1078	APMADTTPNGPQGAGAVQFMMTNKLDTAMWL SRLFTVYCSALFVLPLLGLHEAASFYQRALLANA LTSALRLHQRLPHFQLSRAFLAQALLEDSCHYLL YSLIFVNSYPVTMSIFPVLLFSLLHAATYTKKVL\ DARG\SNSLPLLR\SVLDKLSANQQNILKFIACNEI FLMPATVFMLFSGQGSLLQPFIYYRFLTLRYSSRR NPYCRTLFNELRIVVEHIIMKPACPLFVRRLCLQS IAFISRLAPTVP
3496	A	3	2867	SSRTREMEEKEILRRQIRLLQGLIDDYKTLHGNAP APGTPAASGWQPPTYHSGRAFSARYPRPSRRGYS SHHGPSWRKKYSLVNRPPGPSDPPADHAVRPLH GARGGQPPVPQQHVLERQVQLSQGQNVVIKVKP PSKSGSASASGAQRGSLEEFEDTPWSDQRPREGE GEPPRGQLQPSRPTRARGTCSVEDPLLVCQKEPG KPRMVKSVGSVGDSPREPRRTVSESVIAVKASFP SSALPPRTGVALGRKLGSHSVASCAPQLLGDRRV DAGHTDQPVPSGSVGGPARPASGPRQAREASLV VTCRTNKFRKNNYKWVAASSKSPRVARRALSPR VAAENVCKASAGMANKVEKPQLIADPEPKPRKP ATSSKPGSAPSKYKWKASSPSASSSSSFRWQSEA GSKDHASQLSPVLSRSPSGD\RPALAHSGLKPLSG ETPLSAYKVKTRTKIIRRRGSTSLPGDKKSGTSPA ATAKSHLSLRRRQALRGKSSPVLKKTPNKGLVQ

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SEQ ID NO:	Method	Predicted beginning nucleotide location corresponding to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion
·				VTKHRLCRLPPSRAHLPTKEASSLHAVRTAPTSK VIKTRYRIVKKTPASPLSAPPFPLSLPSWRARRLS LSRSLVLNRLRPVASGGGKAQPGSPWWRSKGYR CIGGVLYKVSANKLSKTSGQPSDAGSRPLLRTGR LDPAGSCSRSLASRAVQRSLAIIRQARQRREKRK EYCMYYNRFGRCNRGERCPYIHDPEKVAVCTRF VRGTCKKTDGTCPFSHHVSKEKMPVCSYFLKGI CSNSNCPYSHVYVSRKAEVCSDFLKGYCPLGAK CKKKHTLLCPDFARRGACPRGAQCQLLHRTQKR HSRRAATSPAPGPSDATARSRVSASHGPRKPSAS QRPTRQTPSSAALTAAAVAAPPHCPGGSASPSSS KASSSSSSSSSPPASLDHE\APSLQEAALAAACSN RLCKLPSFISLQSSPSPGAQPRVRAPRAPLTKDSG KPLHIKPRL
3497	A	1586	141	ATARDLGCARRIDRVVMESTPSRGLNRVHLQCR NLQEFLGGLSPGVLDRLYGHPATCLAVFRELPSL AKNWVMRMLFLEQPLPQAAVALWVKKEFSKA QEESTGLLSGLRIWHTQLLPGGLQGLILNPIFRQN LRIALLGGGKAWSDDTSQLGPDKHARDVPSLDK YAEERWEVVLHFMVGSPSAAVSQDLAQLLSQA GLMKSTEPGEPPCITSAGFQFLLLDTPAQLWYFM LQYLQTAQSRGMDLVEILSFLFQLSFSTLGKDYS VEGMSDSLLNFLQHLREFGLVFQRKRKSRRYYP T/RALAINLSSGVSGAGGTVHQPGFIV\VETNYRL YAYTESELQIALIALFSEMLYPFP\NMVV\ARVTR\ ESVQQAIASGITAQQIIHFLRTRAHPVMLKQTPVL PPTITDQIRLWELERDRLRFTEGVLYNQFLSQVDF ELL\LAHAPKLGVLVFE/NTPAKRLMVVTPAGHS DVKRFWKRQKHSS
3498	A	790		RDLGPAALMTASASSFSSSQGVQQPSIYSFSQITR SLFLSNGVAANDKLLLSSNRITAIVNASVGSGQRI LRG\LQYIKVPVTDARDSRLYDFFDPIADLIHTVS MRQGRTLLNCMAG\MSRSASLCLAYLMKYHSM S\LLDAHTWA/TKSRRPIIRPNNGFWEQLINYEFK LFNNNTVRMINSPVGNIPDIYEKDLRMMISM
3499	A	31		TAGFLLAPLEMQRLLTPVKRILQLTRAVQETSLT PARLLPVAHQRFSTASAVPLAKTDTWPKDVGIL ALEVYFPAQYVDQTDLEKYNNVEAGKYTVGLG QTRMGFCSVQEDINSLCLTVVQRLMERIQLPWD SVGRLEVGTETIIDKSKAVKTVLMELFQDSGNTD IEGIDTTNACYGGTASLFNAANWMESSSWDGRY AMVVCGDIAVYPSGNARPTGGAGAVAMLIGPK APLALERGLRGTHMENVYDFYKPNLASEYPIVD GKLSIQCYLRALDRCYTSYRKKIQNQWKQAGSD RPFTLDDLQYMIFHTPFCKMVQKSLARLMFNDF LSASSDTQTSLYKGLEAFGGLKLEDTYTNKDLD KALLKASQDMFDKKTKASLYLSTHNGNMYTSSL YGCLASLLSHHSAQELAGSRIGAFSYGSGLAASF FSFRVSQDAAPGSPL\DKLVSSTSDLPKRLASRKC VSPEEFTEIMNQREQFYHKVNFSPPGDTNSLFPGT WYLERVDEQHRRKYARRPV
3500	A	185	2692	MLPTEVPQSHPGPSALLLLQLLLPPTSAFFPNIWS LLAAPGSITHQDLTEEAALNVTLQLFLEQPPPGRP PLRLEDFLGRTLLADDLFAAYFGPGSSRRFRAAL GEVSRANAAQDFLPTSRNDPDLHFDAERLGQGR

·				LA - La - La - La - La - La - La - La -
SEQ ID NO:	Method	Predicted beginning	Predicted end nucleotide	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine,
110.		nucleotide	location	I=Isoleucine, K=Lysine, L=Leucine, M=Methionine,
		location	corresponding	N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine,
		corresponding to first amino	to last amino acid residue of	T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion,
		acid residue of	peptide	\=possible nucleotide insertion
	Ì	peptide	sequence	
	ļ	sequence		ARLVGALRETVVAARALDHTLARQRLGAALHA
1				LQDFYSHSNWVELGEQQPHPHLLWPRQELQNLA
}				QVADPTCSDCEELSCPRNWLGFTLLTSGYFGTHP
				PKPPGKCSHGGHFDRSSSQPPRGGINKDSTSPGFS
				PHHMLHLQAAKLALLASIQAFSLLRSRLGDRDFS
ļ	1	İ		RLLDITPASSLSFVLDTTGSMGEEINAAKIQARHL
	}			VEQRRGSPMEPVHYVLVPFHDPGFGPVFTTSDPD
ļ		1	1	SFWQQLNEIHALGGGDEPEMCLSALQLALLHTPP
			ļ	LSDIFVFTDASPKDAFLTNQVESLTQERRCRVTFL
				VTEDTSRVQGRARREILSPLRFEPYKAVALASGG
				EVIFTKDQHIRDVAAIVGESMAALVTLPLDPPVV
	ĺ		[VPGQPLVFSVDGLLQKITVRIHGDISSFWIKNPAG
			ļ	VSQGQEEGGGPLGHTRRFGQFWMVTMDDPPQT
	1			GTWEIQVTAEDTPGVRVQAQTSLDFLFHFGIPME
			ŀ	DGPHPGLYPLTQPVAGLQTQLLVEVTGLGSRAN
				PGDPQPHFSHVILRGVPEGAELGQVPLEPVGPPE
ļ		ļ		RGLLAASLSPTLLSTPRPFSLELIGQDAAGRRLHR
				AAPQPSTVVPVLLELSGPSGFLAPGSKVPLSLRIA SFSGPQDLDLRTFVNPSFSLTSNLSRAHLELNESA
			1	WGRLWLEVPDSAAPDSVVMVTVTAGGREANPV
] ·	PPTHAFLRLLVSAPAPQDRH
3501	A	1245	5815	RRAHPSHSRLSPYLSVSRDPYFFVTVSRTILTLSA
3501		12.5	5015	PAPPRRTPAPSMGTALLQRGGCFLLCLSLLLLGC
				WAELGSGLEFPGAEGQWTRFPKWNACCESEMSF
				QLKTRSARGLVLYFDDEGFCDFLELILTRGGRLQ
				LSFSIFCAEPATLLADTPVNDGAWHSVRIRRQFR
				NTTLFIDQVEAKWVEVKSKRRDMTVFSGLFVGG
Ì	ł		}	LPPELRAAALKLTLASVREREPFKGWIRDVRVNS
			1	SQVLPVDSGEVKLDDEPPNSGGG\SPCEAGEEGE
ł			ĺ	GGVCLNGGVCSVVDDQAVCDCSRTGFRGKDCS QEDNNVEGLAHLMMGDQGKEEYIATFKGSEYF
				CYDLSQNPIQSSSDEITLSFKTLQRNGLMLHTGKS
				ADYVNLALKNGAVSLVINLGSGAFEALVEPVNG
1	ļ		1	KFNDNAWHDVKVTRNLRQHSGIGHAMVTISVD
				GILTTTGYTQEDYTMLGSDDFFYVGGSPSTADLP
				GSPVSNNFMGCLKEVVYKNNDVRLELSRLAKQ
ļ	ļ			GDPKMKIHGVVAFKCENVATLDPITFETPESFISL
				PKWNAKKTGSISFDFRTTEPNGLILFSHGKPRHQ
		ļ		KDAKHPQMIKVDFFAIEMLDGHLYLLLDMGSGT
	 			IKIKALLKKVNDGEWYHVDFQRDGRSGTISVNT
				LRTPYTAPGESEILDLDDELYLGGLPENKAGLVF
			<u> </u>	PTEVWTALLNYGYVGCIRDLFIDGQSKDIRQMA
				EVQSTAGVKPSCSKETAKPCLSNPCKNNGMCRD
				GWNRYVCDCSGTGYLGRSCEREATVLSYDGSM FMKIQLPVVMHTEAEDVSLRFRSQRAYGILMAT
			1	TSRDSADTLRLELDAGRVKLTVNLDCIRINCNSS
			ĺ	KGPETLFAGYNLNDNEWHTVRVVRRGKSLKLT
				VDDQQAMTGQMAGDHTRLEFHNIETGIITERRY
				LSSVPSNFIGHLQSLTFNGMAYIDLCKNGDIDYC
	}	}	}	ELNARFGFRNIIADPVTFKTKSSYVALATLQAYT
				SMHLFFQFKTTSLDGLILYNSGDGNDFIVVELVK
				GYLHYVFDLGNGANLIKGSSNKPLNDNQWHNV
				MISRDTSNLHTVKIDTKITTQITAGARNLDLKSDL
			1	YIGGVAKETYKSLPKLVHAKEGFQGCLASVDLN
<u> </u>		l	<u></u>	G\RLP\DLISDGSFSCNGTDSRRGMWKGPSTT\CQ

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SEQ ID NO:	Method	Predicted beginning nucleotide location corresponding to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion
			* -	EDSCSNQGVCLQQWDGFSCDCSMTSFSGPLCND PGTTYIFSKGGGQITYKWPPNDRPSTRADRLAIGF STVQKEAVLVRVDSSSGLGDYLELHIHQGKIGVK FNVGTDDIAIEESNAIINDGKYHVVRFTRSGGNA TLQVDSWPVIERYPAGRQLTIFNSQATIIIGGKEQ GQPFQGQLSGLYYNGLKVLNMAAENDANIAIVG NVRLVGEVPSSMTTESTATAMQSEMSTSIMETTT TLATSTARRGKPPTKEPISQTTDDILVASAECPSD DEDIDPCEPSSGGLANPTRAGGREPYPGSAEVIRE SSSTTGMVVGIVAAAALCILILLYAMYKYRNRDE GSYHVDESRNYISNSAQSNGAVVKEKQPSSAKSS NKNKKNKDKEYYV
3502	A	394	72	KPAHLPFTVIIMPKRKPSEGAMSDKVKA/KFELQ RRSAGLFSKPTPPKPETRPKKDPANQRQKLPKVR KGKADA/SKEGNSPAEERCSMVQTQKVEGWRSG SELPVALSF
3503	A	43	3358	SGGRGPVRVRSEQLSPSAEQVSQISQISLGRRPLS SLPPPPSRALAPTRAPDTALTIMEVAEVESPLNPS CKIMTFRPSMEEFREFNKYLAYMESKGAHRAGL AKVIPPKEWKPRQCYDDIDNLLIPAPIQQMVTGQ SGLFTQYNIQKKAMTVKEFRQLANSGKYCTPRY LDYEDLERKYWKNLTFVAPIYGADINGSIYDEGV DEWNIARLNTVLDVVEEECGISIEGVNTPYLYFG MWKTTFAWHTEDMDLYSINYLHFGEPKSWYAIP PEHGKRLERLAQGFFPSSSQGCDAFLRHKMTLIS PSVLKKYGIPFDKITQEAGEFMITFPYGYHAGFN HGFNCAESTNFATVRWIDYGKVAKLCTCRKDM VKISMDIFVRKFQPDRYQLWKQGKDIYTIDHTKP TPASTPEVKAWLQRRRKVRKASRSFQCARSTSK RPKADEEEEVSDEVDGAEVPNPDSVTDDLKVSE KSEAAVKLRNTEASSEEESSASRMQVEQNLSDHI KLSGNSCLSTSVTEDIKTEDDKAYAYRSVPSISSE ADDSIPLSTGYEKPEKSDPSELSWPKSPESCSSVA ESNGVLTEGEESDVESHGNGLEPGEIPAVPSGER NSFKVPSIAEGENKTSKSWRHPLSRPPARSPMTL VKQQAPSDEELPEVLSIEEEVEETESWAKPLIHL WQTKPPNFAAEQEYNATVARMKPHCAICTLLMP YHKPDSSNEENDARWETKLDEVVTSEGKTKPLIP EMCFIYSEENIEYSPPNAFLEEDGTSLLISCAKCC VRVHASCYGIPSHEICDGWLCARCKRNAWTAEC CLCNLRGGALKQTKNNKWAHVMCAVAVPEVR FTNVPERTQIDVGRIPLQRLKLKCIFCRHRVKRVS GACIQCSYGRCPASFHVTCAHAAGVL\MEPDDW PYVVNITCFRHKVNPNVKSKACEKVISVGQTVIT KHRNTRYYSCRVMAVTSQTFYEVMFDDGSFSRD TFPEDIVSRDCLKLGPPAEGEVVQVKWPDGKLY GAKYFGSNIAHMYQVEFEDGSQIAMKREDIYTL DEELPKRVKARFVSAGRCHLGTCQVNSLSSPHVS QAQQETYLGFWINSKKSQCNIFLSGTY
3504	A	1124	139	RGEEQFDAEFRRFACLGFGERLQEFSRLLRAVHR SRAWTCYLAIRMLMATCCPSPTTTACTGPWQRA PPLRLLVQKREADSSGLAFASNSLQRRKKGLLLR PVAPLRTRPPLLISLPQDFRQVSSVIDVDLLPETH RRVRLHKHGSDRPLGFYIRDGMSVRVAPQG\LER VPGIFISRLVRGGLAESTGLLAVSDEILEVNGIEV

AGKTLNOVTDMMVANSHNLIVTVKPANQRNN VVRGASGRLTGPPSAGROPAEPDSDDDSDLVI NROPPSSNGLSOGPPCVDLHPGCRHPGTRSSLPS LDDQSQASSWGSRIRGDGSGFSL LDDQSQASSWGSRIRGDGSGFSL LDDQSQASSWGSRIRGDGSGFSL LDDQSQASSWGSRIRGDGSGFSL LDDQSQASSWGSRIRGDSGFSL LDDQSQASSWGSRIRGDGSGFSL LDDQSQASSWGSRIRGDGSGFSL SALCPKFLHTNSTSHTWPFSAVAELLDNAYDPD NAKQIVIDKTVINDHICLTFTDNGNGMTSDKLH KMLSFGFSDKVTMNGHPVPGLYGNGFKSGSMI LGKDAIVFIKNGESMSVGLLSQTYLEVIKABI LGKDAIVFIKNGESMSVGLLSQTYLEVIKABI LGKDAIVFIKNGESMSVGLLSQTYLEVIKABI LGKDAIVFIKNGESMSVGLLSQTYLEVIKABI LGKDAIVFIKNGESMSVGLLSQTYLEVIKABI LGKDAIVFIKNGESMSVGLLSQTYLEVIKABI LGKDAIVFIKNGESKSSVLABLEHSLFST QKLLAELDAIIGKKGTRIIIWNLRSYKNATEDFS KDKYMRIPEDLDEITGKKGYKKQERMDQIAPE DYSLRAYCSILYLKPRNOJILRGQKVKTQLVSKS LAYERDVYRFKFLSKTVRITGFFORCNKOHYD KDYLRAYCSILYLKPRNOJILRGQKVKTQLVSKS LAYERDVYRFKFLSKTVRITGFFORCNKOHYD MMYHRNRLIKAYEKVGCQLRANNMGVGVVQI ECNFLKPTHNKQDFDYTNEYRLTTIALGEKLND YWNEMKVKKNTEYPLIN,PVEDIQKRPDQTWV CDACLKWRKLPDGMDQLPEKWYCSNNPDPQR NCEVPEEPEDEDLVHPTYEELTYKKTNKEKFRIR PEMBRINABLLFRITALSTPSYSSPKESVSKRR LSEGTNSYATRLLNNHQVPPQSEPESNSLKRR LSEGTNSYATRLLNNHQVPPQSEPESNSLKRR LSEGTNSYATRLLNNHQVPPQSEPESNSLKRR LENSTINANRRLSSQFENSYVKGIDDDEDVI LEENSTFKPAVDHDIDMKSEQSHVEQGGVQVEI UVGKSEFCGGTGSTSTSSSRCQGGTNCTATQTEVP LVVKKEETVEDEEDVRNDAVILPSCVEABAKHT TQETTDKSADDAGCQLQELRQCLLVTEEKEN KRQCHMFTDQIKVLQQRILEMNDKYVKKETCH QSTETDAVTLLESINGKSSEPDHMVSQVQQALE IERLKKQCSALQHVKABCCSQCSNNESKSEMDEN AVQLDDVFRQLDKCSERDQVKSSVELLEMENS SVHIMDGGSLKLRSLRVNVQQLLAMIVPDLDL QVNYDVDVDELIGQVVEQMSEISST GRERAFAGGATAGSGRGSFTGPAGG GRTRRAMDRPAAAAAACCEGGGGFRGPAGAG RPPRAAGGATAGSGRGSVTSENVSRDYKARRDGNKLA QMEEJALFFESIKAYKDVMYCPFMGAVSC QSHGDNSCGIENCKDMRNRLAYKQEQSKLU IFENLINKHAFPLSNGQALAFSYKEKPINGWK YDPVSEYKRQGLPNESWKISKINSNYCEPCDTYP, IUVPTSVKDDDLSKVAVFLAKGGRYVSKSINGN PPSAFTTSVGPLVTPSVKDGLSKVAVFLAKGRYVVLSWIPI SQATTITCSQPLVQPNDKRCKEDEKVLQTIMDA AQSHKLIIFDARQNSVADTNKTKGGGYESSEAY NAELVFLEINHIPMRSLSKLKERVYPSIDBAR LSNYDGTHWLEYRIMLLAGAVRIADKSTKIG TLYEKSWIFGGJAFRALEWOTHADADADRSP LQVPQCVWQMTRQPFSAFFRILLITILLBUT VYHCSGWWBTANQLTSLAWALDSNYTKTIG TLYEKSWIFGGGRFALERVOTHADADADRSP LQVPQCVWQMTRQPFSAFFRILL	SEQ ID NO:	Method	Predicted beginning nucleotide location corresponding to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion
SALCPKFLHTNSTSHTWPFSAVABLIDNAYDPDV NAKQIWIDKTVINDHICLTFTDNGNGMISDKLH KMLSFGFSIKVTMNGHVPVGLYGNGFKSGSMI LGRDAIVFTKNGESMSVGLLSQTYJLEVIKAEHV VVPIVAFNKHRQMINLAESKASLAALHEISLFST QKLLAELDAIIGKKGTRIIIWNLRSYKNATEFDFI KDKYDIRIPEDLDEITGKKGYKKQERMDQIAPE DYSLRAYCSIL YLRPRMQHILAGGKVKTQLVSKS LAYERDVYRPKFLSKTVRITFGFNCRNKDHYGI MMYHRNRLIKAYEKVGCQLRANNMGVGVVGI ECNFLKPHINKQDFDVTNEVRLITITALGEKLND YWNEMKVKKNTEYPLNLPVEDIGKRPDQTWV CDACLKWRKLPDGMDQLPKWYCSNNPDPQF NCEVPEEPEDEDLVHPTYEKTYKKTNKEKFRIR PEMIPRINAELLFRIPTALSTPSYSSPKESVSKRR LSEGTINSYATRLINNHQVPPQSEPESNSLKRRLS TRSSILNAKNRRLISSQFENSVYKGDDDDEDVI LEENSTPKPAVDHDDMKSEQSHVEGGQVQVEI VGDSEPCGGTGSTSTSSSRCDQGNTAATQTEVP LVVKKEETVEDEIDVRNDAVILPSCVEAEAKHH TQRTTDKSADDAGGQLQELRNQLLLVTEEKEN KRQCHMFTDQIKVLQQRILEMNDKYVKKETCH QSTETDAVFLLESINGKSESPDHMVSQVQALE IERKKQCSALQHYKABCSQCSNNESKSEMDE AVQLDDVFRQLDKCSIERDQYKSEVELLEMEKS QIRSQCEBLKTEVFQLKSTNQQTATDVSTSSNIE SVNHMDGESLKLRSLRVNVGQLLAMIVPDLDL QVNYDVDVDELIGQVVEQMSEISST SYNHMDGESLKLRSLRVNVGQLLAMIVPDLDL QVNYDVDVDELIGQVVEQMSEISST TYTDFKLYFKAGGRRQAAKPSRPPLPSRRRLPQ GRTRRAMDRPAAAAAGCEGGGPNPGPAGG RPPRAAGGATAGSRQPSVETLDSPTGSHVEWCK QLIAATISSQISGSVTSENVSRDYKALRDGNKLA QMEEAPLFPGESIKAIVKDVMYICPFMGAVSGTI TYTDFKLYFKNVERDPHIEDJVPLGVISRVEKIG QSHGDNSCGEIVCKDMRNLRLAYKQEEQSKL IFENLNKHAFPLSNQALFAFSYKEKFPINGWK' YDDYSEYKRQGLPNESWKISKINSNYEPCDTYP- IIVYPTSVKDDDLSKVAVFLAKGRYPVLSWHPP SQATITRCSQPLVGPNDKRCKEDEKYLQTIMDA AQSHKLIFDARQNSVADTNKTRGGGYESESAY NAELVFLEHNIHVMRESLRKLKEIVYPSIDEAR LSNVDGTHWLFYRMLLAGAVRIADKEGGKTS VVHCSDGWDRTAQLTSLAMLMLDSYYRTIKGF TLVEKEWISFGRFFALRVGHGNDNHADADARSP LQPVDCVWQMTGQFFSAFFENFELITLIDHLYS CLEGTFTLCNCEQQFKEDVYTKTISLWSYNSQI DEFSNFFFVNYENHVLYPVASLSLHLEWNYNY RWNPRMRQMPHHONLKELLAVRAELQKRVE	·				VVRGASGRLTGPPSAGPGPAEPDSDDDSSDLVIE NRQPPSSNGLSQGPPCWDLHPGCRHPGTRSSLPS LDDQEQASSGWGSRIRGDGSGFSL
3506 A 2 2120 RPPEAGGRYRAGGRRQAAKPSRPPLPSRRRLPQ GRTRRAMDRPAAAAAAGCEGGGGPNPGPAGG RPPRAAGGATAGSRQPSVETLDSPTGSHVEWCK QLIAATISSQISGSVTSENVSRDYKALRDGNKLA QMEEAPLFPGESIKAIVKDVMYICPFMGAVSGTI TVTDFKLYFKNVERDPHFILDVPLGVISRVEKIG QSHGDNSCGIEIVCKDMRNLRLAYK\QEEQSKL IFENLNKHAFPLSNGQALFAFSYKEKFPINGWKV YDPVSEYKRQGLPNESWKISKINSNYEFCDTYP- IIVVPTSVKDDDLSKVAVFLAKGRVPVLSWIHPP SQATITRCSQPLVGPNDKRCKEDEKYLQTIMDA AQSHKLIIFDARQNSVADTNKTKGGGYESESAY NAELVFLEIHNIHVMRESLRKLKEIVYPSIDEAR* LSNVDGTHWLEYIRMLLAGAVRIADKIESGKTS VVHCSDGWDRTAQLTSLAMLMLDSYYRTIKGF TLVEKEWISFGHRFALRVGHGNDNHADADRSP- LQFVDCVWQMTRQFPSAFEFNELFLITILDHLYS CLFGTFLCNCEQQRFKEDVYTKTISLWSYINSQI DEFSNPFFVNYENHVLYPVASLSHLELWVNYYV RWNPRMRPQMPIHQNLKELLAVRAELQKRVEG	3505	A	3	2898	SCRSATSQSGCGGGRSWLCSSLKMAAQPPRGIRL SALCPKFLHTNSTSHTWPFSAVAELIDNAYDPDV NAKQIWIDKTVINDHICLTFTDNGNGMTSDKLH KMLSFGFSDKVTMNGHVPVGLYGNGFKSGSM/R LGKDAIVFTKNGESMSVGLLSQTYLÆVIKAEHV VVPIVAFNKHRQMINLAESKASLAAILEHSLFSTE QKLLAELDAIIGKKGTRIIIWNLRSYKNATEFDFE KDKYDIRIPEDLDEITGKKGYKKQERMDQIAPES DYSLRAYCSILYLKPRMQIILRGQKVKTQLVSKS LAYIERDVYRPKFLSKTVRITFGFNCRNKDHYGI MMYHRNRLIKAYEKVGCQLRANNMGVGVVGII ECNFLKPTHNKQDFDYTNEYRLTITALGEKLND YWNEMKVKKNTEYPLNLPVEDIQKRPDQTWVQ CDACLKWRKLPDGMDQLPEKWYCSNNP\DPQFR NCEVPEEPEDEDLVHPTYEKTYKKTNKEKFRIRQ PEMIPRINAELLFRPT\ALSTPS\FSSPKESVSKR/RH LSEGTNSYATRLLNNHQVPPQSEPESNSLKRRLS TRSSILNAKNRRL\SSQF\ENSVYKG\DDDDEDVII LEENSTPKPAVDHDIDMKSEQSHVEQGGVQVEF VGDSEPCGQTGSTSTSSSRCDQGNTAATQTEVPS LVVKKEETVEDEIDVRNDAVILPSCVEAEAKIHE TQETTDKSADDAGCQLQELRNQLLLVTEEKENY KRQCHMFTDQIKVLQQRILEMNDKYVKKETCH QSTETDAVFLLESINGKSESPDHMVSQYQQALEE IERLKKQCSALQHVKAECSQCSNNESKSEMDEM AVQLDDVFRQLDKCSIERDQYKSEVELLEMEKS QIRSQCEELKTEVEQLKSTNQQTATDVSTSSNIEE SVNHMDGESLKLRSLRVNVGQLLAMIVPDLDLQ OVNYDVDVDEILGOVVEQMSEISST
LQREVATRAVSSSSERGSSPSHFATSVHTLV 3507 A 1 2169 GSSIKIRLTVLCAKNLAKKDFFRLPDPF\AKIVVL		-			RPPEAGGRYRAGGRRQAAKPSRPPLPSRRRLPQG GRTRRAMDRPAAAAAAGCEGGGGPNPGPAGGR RPPRAAGGATAGSRQPSVETLDSPTGSHVEWCK QLIAATISSQISGSVTSENVSRDYKALRDGNKLA QMEEAPLFPGESIKAIVKDVMYICPFMGAVSGTL TVTDFKLYFKNVERDPHFILDVPLGVISRVEKIGA QSHGDNSCGIEIVCKDMRNLRLAYK\QEEQSKLG IFENLNKHAFPLSNGQALFAFSYKEKFPINGWKV YDPVSEYKRQGLPNESWKISKINSNYEFCDTYPA IIVVPTSVKDDDLSKVAVFLAKGRVPVLSWIHPE SQATITRCSQPLVGPNDKRCKEDEKYLQTIMDAN AQSHKLIIFDARQNSVADTNKTKGGGYESESAYP NAELVFLEIHNIHVMRESLRKLKEIVYPSIDEARW- LSNVDGTHWLEYIRMLLAGAVRIADKIESGKTSV VVHCSDGWDRTAQLTSLAMLMLDSYYRTIKGFE TLVEKEWISFGHRFALRVGHGNDNHADADRSPIF LQFVDCVWQMTRQFPSAFEFNELFLITILDHLYS CLFGTFLCNCEQQRFKEDVYTKTISLWSYINSQL DEFSNPFFVNYENHVLYPVASLSHLELWVNYYV RWNPRMRPQMPIHQNLKELLAVRAELQKRVEG

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				GSGQCHSTDTVKNTLDPKWNQHYDLYVGKTDSI TISVWNHKKIHKKQGAGFLGCVRLLSNAISRLKD TGYQRLDLCKLNPSDTDAVRGQIVVSLQTRDRIG TGGSVVDCRGLLENEGTVYEDSGPGRPLSCFME EPAPYTDSTGAAAGGGNCRFVESPSQDQRLQAQ RLRNPDVRGSLQTPQNRPHGHQSPELPEGYEQRT TVQGQVYFLHTQTGVSTWHDPRIPRDLNSVNCD ELGPLPPGWEVRSTVSGRIYFVDHNNRTTQFTDP RLHHIMNHQCQLKEPSQPLPLPSEGSLEDEELPA QRYERDLVQKLKVLRHELSLQQPQAGHCRIEVS REEIFEESYRQIMKMRPKDLKKRLMVKFRGEEG LDYGGVAREWLYLLCHEMLNPYYGLFQYSTDNI YMLQINPDSSINPDHLSYFHFVGRIMGLAVFHGH YINGGFTVPFYKQLLGKPIQLSDLESVDPELHKSL VWILENDITPVLDHTFCVEHNAFGRILQHELKPN G\RNVPVTEENKKEYVRLYVNWRFMRGIEAQFL ALQKGFNELIPQHLLKPFDQKELELIIGGLDKIDL NDWKSNTRLKHCVADSNIVRWFWQAVETFDEE RRARLLQFVTGSTRVPLQGFKALQGSTG\AAGPR LFTIHLIDANTDNLRKAHTCFNRIDIPPYESYEKL
3508	A	3	6388	ILYINPADLGWNPPVSSWIEKREIQTERANLTILF DKYLPTCLDTLRTRFKKIIPIPEQSMVQMVCHLLE CLLTTEDIPADCPKEIYEHYFVFAAIWAFGGAMV QDQLVDYRAEFSKWWLTEFKTVKFPSQGTIFDY YIDPETKKFEPWSKLVPQFEFDPEMPLQACLVHT SETIRVCYFMERLMARQRPVMLVGTAGTGKSVL VGAKLASLDPEAYLVKNVPFNYYTTSAMLQAVL EKPLEKKAGRNYGPPGNKKLIYFIDDMNMPEVD AYGTVQPHTIIRQHLDYGHWYDRSKLSLKEITNV QYVSCMNPTAGSFTINPRLQRHFSVFVLSFPGAD ALSSIYSIILTQHLKLGNFPASLQKSIPPLIDLALAF HQKIATTFLPTGIKFHYIFNLRDFANIFQGILFSSV ECVKSTWDLIRLYLHESNRVYRDKMVEEKDFDL FDKIQTEVLKKTFDDIEDPVEQTQSPNLYCHFAN GIGEPKYMPVQSWELLTQTLVEALENHNEVNTV MDLVLFEDAMRHVCHINRILESPRGNALLVGVG GSGKQSLTRLAAFISSMDVFQITLRKGYQIQDFK MDLASLCLKAGVKNLNTVFLMTDAQVADERFL VLINDLLASGEIPDLYSDDEVENIISNVRNEVKSQ GLVDNRENCWKFFIDRIRRQLKVTLCFSPVGNKL RVRSRKFPAIVNCTAIHWFHEWPQQALESVSLRF LQNTEGIEPTVKQSISKFMAFVHTSVNQTSQSYLS NEQRYNYTTPKSFLEFIRLYQSLLHRHRKELKCK TERLENGLLKLHSTSAQVDDLKAKLAAQEVELK QKNEDADKLIQVVGVETDKVSREKAMADEEEQ KVAVIMLEVKQKQKDCEEDLAKAEPALTAAQA ALNTLNKTNLTELKSFGSPPLAVSNVSAAVMVL MAPRGRVPKDRSWKAAKVTMAKVDGFLDSLIN FNKENIHENCLKAIRPYLQDPEFNPEFVATKSYA AAGLCSWVINIVRFYEVFCDVEPKRQALNKATA DLTAAQEKLAAIKAKIAHLNENLAKLTARFEKA TADKLKCQQEAEVTAVTISLANRLVGGLASENV RWADAVQNFKQQERTLCGDILLITAFISYLGFFT KKYRQSLLDRTWRPYLSQLKTPIPVTPALDPLRM

-075 C T		D - 3° 4-3	D22-4-3	Amino cold commons (A - Alentes Co-Constant W. Account A. 17
SEQ ID NO:	Method	Predicted beginning nucleotide location corresponding to first amino	Predicted end nucleotide location corresponding to last amino acid residue of	Amino acid sequence (A=Alauine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion,
		acid residue of peptide sequence	peptide sequence	├─possible nucleotide insertion
			-	LMDDADVAAWQNEGLPADRMSVENATILINCE
				RWPLMVDPQLQGIKWIKNKYGEDLRVTQIGQKG
		r		YLQIIEQALEAGAVVLIENLEESIDPVLGPLLGRE VIKKGRFIKIGDKECEYNPKFRLILHTKLANPHYQ
				PELQAQATLINFTVTRDGLEDQLLAAVVSMERP
				DLEQLKSDLTKQQNGFKITLKTLEDSLLSRLSSAS
				GNFLGETVLVENLEITKQTAAEVEKKVQEAKVT
		1		EVKINEAREHYRPAAARASLLYFIMNDLSKIHPM
				YQFSLKAFSIVFQKAVERAAPDESLRERVANLID
			ļ	SITFSVYQYTIRGLFECDKLTYLAQLTFQILLMNR EVNAVELDFLLRSPVQTGTASPVEFLSHQAWGA
				VKVLSSMEEFSNLDRDIEGSAKSWKKFVESECPE
				KEKLPQEWKNKTALQRLCMLRAMRPDRMTYAL
				RDFVEEKLGSKYVVGRALDFATSFEESGPATPMF
				FILSPGVDPLKDVESQGRKLGYTFNNQNFHNVSL
				GQGQEVVAEAALDLAAKKGHWVILQNTLEMCS RETEFKSILFALCYFHAVVAERRKFGPQGWNRSY
				PFNTGDLTISVNVLYNFLEANAKVPYDDLRYLFG
				EIMYGGHITDDWDRRLCRTYLGEFIRPEMLEGEL
				SLAPGFPLPGNMDYNGYHQYIDAELPPESPYLYG
				LHPNAEIGFLTQTSEKLFRTVLELQPRDSQARDG
		1		AGATREEKVKALLEEILERVTDEFNIPELMAKVE ERTPYIVVAFQECGRMNILTREIQRSLRELELGLK
				GELTMTSHMENLQNALYFDMVPESWARRAYPS
				TAGLAAWFPDLLNRIKELEAWTGDFTMPSTVWL
		ļ		TGFFNPQSFLTAIMQSTARKNEWPLDQMALQCD
				MTKKNREEFRSPPREGAYIHGLFMEGACWDTQA
				GIITEAKLKDLTPPMPVMFIKAIPAD\RQDCGHVY SCPVTKTSQ\RDPTYVWTFNLKTKENPSKWVLA
				GVALLLQI
3509	A	3	6388	ILYINPADLGWNPPVSSWIEKREIQTERANLTILF
				DKYLPTCLDTLRTRFKKIIPIPEQSMVQMVCHLLE
				CLLTTEDIPADCPKEIYEHYFVFAAIWAFGGAMV QDQLVDYRAEFSKWWLTEFKTVKFPSQGTIFDY
1				YIDPETKKFEPWSKLVPQFEFDPEMPLQACLVHT
		}		SETIRVCYFMERLMARQRPVMLVGTAGTGKSVL
;	'			VGAKLASLDPEAYLVKNVPFNYYTTSAMLQAVL
				EKPLEKKAGRNYGPPGNKKLIYFIDDMNMPEVD
				AYGTVQPHTIIRQHLDYGHWYDRSKLSLKEITNV QYVSCMNPTAGSFTINPRLQRHFSVFVLSFPGAD
				ALSSIYSIILTQHLKLGNFPASLQKSIPPLIDLALAF
				HQKIATTFLPTGIKFHYIFNLRDFANIFQGILFSSV
				ECVKSTWDLIRLYLHESNRVYRDKMVEEKDFDL
				FDKIQTEVLKKTFDDIEDPVEQTQSPNLYCHFAN
				GIGEPKYMPVQSWELLTQTLVEALENHNEVNTV MDLVLFEDAMRHVCHINRILESPRGNALLVGVG
		1		GSGKQSLTRLAAFISSMDVFQITLRKGYQIQDFK
				MDLASLCLKAGVKNLNTVFLMTDAQVADERFL
				VLINDLLASGEIPDLYSDDEVENIISNVRNEVKSQ
		1	Ì	GLVDNRENCWKFFIDRIRRQLKVTLCFSPVGNKL
		}		RVRSRKFPAIVNCTAIHWFHEWPQQALESVSLRF LQNTEGIEPTVKQSISKFMAFVHTSVNQTSQSYLS
				NEORYNYTTPKSFLEFIRLYQSLLHRHRKELKCK
		}		TERLENGLLKLHSTSAQVDDLKAKLAAQEVELK
L	L		L	QKNEDADKLIQVVGVETDKVSREKAMADEEEQ

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				KVAVIMLEVKQKQKDCEEDLAKAEPALTAAQA ALNTLNKTNLTELKSFGSPPLAVSNVSAAVMVL MAPRGRVPKDRSWKAAKVTMAKVDGFLDSLIN FNKENIHENCLKAIRPYLQDPEFNPEFVATKSYA AAGLCSWVINIVRFYEVFCDVEPKRQALNKATA DLTAAQEKLAAIKAKIAHLNENLAKLTARFEKA TADKLKCQQEAEVTAVTISLANRLVGGLASENV RWADAVQNFKQQERTLCGDILLITAFISYLGFFT KKYRQSLLDRTWRPYLSQLKTPIPVTPALDPLRM LMDDADVAAWQNEGLPADRMSVENATILINCE RWPLMVDPQLQGIKWIKNKYGEDLRVTQIGQKG YLQIIEQALEAGAVVLIENLEESIDPVLGPLLGRE VIKKGRFIKIGDKECEYNPKFRLILHTKLANPHYQ PELQAQATLINFTVTRDGLEDQLLAAVVSMERP DLEQLKSDLTKQQNGFKITLKTLEDSLLSRLSSAS GNFLGETVLVENLEITKQTAAEVEKKVQEAKVT EVKINEAREHYRPAAARASLLYFIMNDLSKIHPM YQFSLKAFSIVFQKAVERAAPDESLRERVANLID SITFSVYQYTIRGLFECDKLTYLAQLTFQILLMNR EVNAVELDFLLRSPVQTGTASPVEFLSHQAWGA VKVLSSMEEFSNLDRDIEGSAKSWKKFVESECPE KEKLPQEWKNKTALQRLCMLRAMRPDRMTYAL RDFVEEKLGSKYVVGRALDFATSFEESGPATPMF FILSPGVDPLKDVESQGRKLGYTFNNQNFHNVSL GQGQEVVAEAALDLAAKKGHWVILQNTLEMCS RETEFKSILFALCYFHAVVAERRKFGPQGWNRSY PFNTGDLTISVNVLYNFLEANAKVPYDDLRYLFG EIMYGGHITDDWDRRLCRTYLGEFIRPEMLEGEL SLAPGFPLPGNMDYNGYHQYIDAELPPESPYLYG LHPNAEIGFLTQTSEKLFRTVLELQPRDSQARDG AGATREEKVKALLEEILERVTDEFNIPELMAKVE ERTPYTVVAFQECGRMNILTREIQRSLRELELGLK GELTMTSHMENLQNALYFDMVPESWARRAYPS TAGLAAWFPDLLNRIKELEAWTGDFTMPSTVWL TGFFNPQSFLTAIMQSTARKNEWPLDQMALQCD MTKKNREEFRSPPREGAYHGLFMEGACWDTQA GITTEAKLKDLTPPMPVMFIKAIPAD\RQDCGHVY SCPVTKTSQ\RDPTYVWTFNLKTKENPSKWVLA GVALLLQI
3510	A	390	3330	AAGSGSRPPAPAARKMADLAECNIKVMCRFRPL NESEVNRGDKYIAKFQGEDTVVIASKPYAFDRVF QSSTSQEQVYNDCAKKIVKDVLEGYNGTIFAYG QTSSGKTHTMEGKLHDPEGMGIIPRIVQDIFNYIY SMDENLEFHIKVSYFEIYLDKIRDLLDVSKTNLSV HEDKNRVPYVKGCTERFVCSPDEVMDTIDEGKS NRHVAVTNMNEHSSRSHSIFLINVKQENTQTEQK LSGKLYLVDLAGSEKVSKTGAEGAVLDEAKNIN KSLSALGNVISALAEGSTYVPYRDSKMTRILQDS LGGNCRTTIVICCSPSSYNESETKSTLLFGQRAKTI KNTVCVNVELTAEQWKKKYEKEKEKNKILRNTI QWLENELNRWRNGETVPIDEQFDKEKANLEAFT VDKDITLTNDKPATAIGVIGNFTDAERRKCEEEIA KLYKQLDDKDEEINQQSQLVEKLKTQMLDQEEL LASTRRDQDNMQAELNRLQAENDASKEEVKEV

SEQ ID NO:	Method	Predicted beginning nucleotide location corresponding to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion
·	·			QKSATLASIDAELQKLKEMTNHQKKRAAEMMA SLLKDLAEIGIAVGNNDVKQPEGTGMIDEEFTVA RLYISKMKSEVKTMVKRCKQLESTQTESNKKME ENEKELAACQLRISQHEAKIKSLTEYLQNVEQKK RQLEESVDALSEELVQLRAQEKVHEMEKEHLNK VQTANEVKQAVEQQIQSHRETHQKQISSLRDEVE AKAKLITDLQDQNQKMMLEQERLRVEHEKLKA TDQEKSRKLHELTVMQDRREQARQDLKGLEETV AKELQTLHNLRKLFVQDLATRVKKSAEIDS\DDT GGSAAQKQKISFLENNLE\QLTKSAQTSWYRDNA DLRCELPKLEKRLRATAERVKALESALKEAKEN ASRDRKRYQQEVDRIKEAVRSKNMARRGHSAQI AKPIRPGQHPAASPTHPSAIRGGGAFVQNSQPVA VRGGGGKQV
3511	A	1	1757	MASVQASRRQWCYLCDLPKMPWAMVWDFSEA VCRGCVNFEGADRIELLIDAARQLKRSHVLPEGR SPGPPALKHPATKDLAAAAAQGPQLPPPQAQPQP SGTGGGVSGQDRYDRATSSGRLPLPSPALEYTLG SRLANGLGREEAVAEGARRALLGSMPGLMPPGL LAAAVSGLGSRGLTLAPGLSPARPLFGSDFEKEK QQRNADCLAELNEAMRGRAEEWHGRPKAVREQ LLALSACAPFNVRFKKDHGLVGRVFAFDATARP PGYEFELKLFTEYPCGSGNVYAGVLAVARQMFH DALREPGKALASSGFKYLEYERRHGSGEWRQLG ELLTDGVRSFREPAPAEALPQQYPEPAPAALCGP PPRAPSRNLAPTPRRRKASPEPEGEAAGKMTTEE QQQRHWVAPGGPYSAETPGVPSPIAALKNVAEA LGHSPKDPGGGGGPVRAGGASPAASSTAQPPTQ HRLVARNGEAEVSPTAGAEAVSGGGSGTGATPG APLC\CTLCRERLEDTHFVQ\CPPVPEHKFCFPCSR KFIKAQGPAGE\VYCPSGDKCPLVGSSVPWAFMQ GEIATILAGDIKVKKERDP
3512	A	3	1994	NTNSSSVTNSAAGVEDLNIVQVTVPDNEKERLSS IEKIKQLREQVNDLFSRKFGEAIGVDFPVKVPYR KITFNPGCVVIDGMPPGVVFKAPGYLEISSMRRIL EAAEFIKFTVIRPLPGLELSNGEYSTVGKRKIDQE GRVFQEKWERAYFFVEVQNISTCLICKRSMSVSK EYNLRHYQTNHSKHYDQYMERMRDEKLHELK KGLRKYLLGLSDTECPEQKQVFANPSPTQKSPVQ PVEDLAGNLWEKLREKIRSFVAYSIAIDEITDINN TTQLAIFIRGVDENFDVSEELLDTVPMTGTKSGN EIFSRVEKSLKNFCINWSKLVSVASTGTPPMVDA NNGLVTKLKSRVATFCKGAELKSICCIIHPESLCA QKLKMDHVMDVVVKSVNWICSRGLNHSEFTTL LYELDSQYGSLLYYTEIKWLSRGLVLKRFFESLE EIDSFMSSRGKPLPQLSSIDWIRDLAFLVDMTMH LNALNISLQGHSQIVTQMYDLIRAFLAKLCLWET HLTRNNLAHFPTLKLVSRNESDGLNYIPKIAELK TEFQKRLSDFKLYESELTLFSSPFSTKIDSVHEELQ MEVIDLQCNTVLKTKYDKVGIPEFYKYLWGSYP KYKHHCAKILSMFGSTYICEQLFSIMKLSKTKYC SQLKDSQWDSVLHIAT
3513	Á	1836	513	FKSLLSVKWFCFSILVLIFLGTRCYWEMTQSRPSP DPHRGRWEGGRSRPKGGEEGRRRTRVPGLVTAS GPGNPLPDRLGEMAGGRHRRVVGTLHLLLLVAA

SEQ ID NO:	Method	Predicted beginning nucleotide location corresponding to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding t last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion
				LPWASRGVSPSASAWPEEKNYHQPAILNSSALRQ IAEGTSISEMWQNDLQPLLIERYPGSPGSYAARQ HIMQRIQRLQADWVLEIDTFLSQTPYGYRSFSNII STLNPTAKRHLVLACHYDSKYFSHW\NNRVFVG ATDSAVPCAMMLELARALDKKLLSLKTVSDSKP DLSLQLIFFDGEEAFLHWSPQDSLYGSRHLAAKM ASTPHPPGARGTSQLHGMDLLVLLDLIGAPNPTF PNFFPNSARWFERLQAIEHELHELGLLKDHSLEG RYFQNYSYGGVIQDDHIPFLRRGVPVLHLIPSPFP EVWHTMDDNEENLDESTIDNLNKILQVFVLEYL HL
3514	A	1836	513	FKSLLSVKWFCFSILVLIFLGTRCYWEMTQSRPSP DPHRGRWEGGRSRPKGGEEGRRRTRVPGLVTAS GPGNPLPDRLGEMAGGRHRRVVGTLHLLLLVAA LPWASRGVSPSASAWPEEKNYHQPAILNSSALRQ IAEGTSISEMWQNDLQPLLIERYPGSPGSYAARQ HIMQRIQRLQADWVLEIDTFLSQTPYGYRSFSNII STLNPTAKRHLVLACHYDSKYFSHW\NNRVFVG ATDSAVPCAMMLELARALDKKLLSLKTVSDSKP DLSLQLIFFDGEEAFLHWSPQDSLYGSRHLAAKM ASTPHPPGARGTSQLHGMDLLVLLDLIGAPNPTF PNFFPNSARWFERLQAIEHELHELGLLKDHSLEG RYFQNYSYGGVIQDDHIPFLRRGVPVLHLIPSPFP EVWHTMDDNEENLDESTIDNLNKILQVFVLEYL HL
3515	A	114	754	LCRDLTTTMSSKRTKTKTKKRPQRATSNVFAMF DQSQIQEFKEAFNMIDQNRDGFIDKEDLHDMLAS LGKNPTDEYLDAMMNEAPGPINFTMFLTMFGEK LNGTDPEDVIRNAFACFDEEATGTIQEDYLRELL TT\MGDRF\TDE\EVDELYREAPI\DKKGGIFNYI\E FTRHLETGGPKDKDDRKITFQIPSPNVPWLATFG VFLEIFLLHGP
3516	A	1	5169	MAAAPSALLLIPPFPVLSTYRLQSRSRPSAPETDD SRVGGIMRGEKNYYFRGAAGDHGSCPTTTSPLA SALLMPSEAVSSSWSESGGGLSGGDEEDTRLLQL LRTARDPSEAFQALQAALPRRGGRLGFPRRKEAL YRALGRVLVEGGSDEKRLCLQLLSDVLRGQGEA GQLEEAFSLALLPQLVVSLREENPALRKDALQIL HICLKRSPGEVLRTLIQQGLESTDARLRASTALLL PILTTEDLLLGLDLTEVIISLARKLGDQETEEESE TAFSALQQIGERLGQDRFQSYISRLPSALRRHYN RRLESQFGSQVPYYLELEASGFPEDPLPCAVTLS NSNLKFGIIPQELHSRLLDQEDYKNRTQAVEELK QVLGKFNPSSTPHSSLVGFISLLYNLLDDSNFKVV HGTLEVLHLLVIRLGEQVQQFLGPVIAASVKVLA DNKLVIKQEYMKIFLKLMKEVGPQQVLCLLLEH LKHKHSRVREEVVNICICSLLTYPSEDFDLPKLSF DLAPALVDSKRRVRQAALEAFAVLASSMGSGKT SILFKAVDTVELQDNGDGVMNAVQARLARKTLP RLTEQGFVEYAVLMPSSAGGRSNHLAHGADTD WLLAGNRTQSAHCHCGDHVRDSMHIYGSYSPTI CTRRVLSAGKGKNKLPWENEQPGIMGENQTSTS KDIEQFSTYDFIPSAKLKLSQGMPVNDDLCFSRK RVSRNLFQNSRDFNPDCLPLCAAGTTGTHQTNLS GKCAQLGFSQICGKTGSVGSDLQFLGTTSSHQEK

SEQ ID NO:	Method	Predicted beginning nucleotide location corresponding to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding t last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion
				ILPSYPVSSPRTSPKHTSPLIISPKKSQDNSVNFSNS WPLKSFEGLSKPKSHRRSLSAQKSS\DPTGR\NHG \ENSQEKPP\VQLTPAL\VRSPSSRRGLNGTKPVPPI P\RGISLLPDKADLSTVGHKKKEPDDIWKCEKDS LPIDLSELNFKDKDLDQEEMHSSLRSLRNSAAKK RAKLSGSTSDLESPDSAMKLDLTMDSPSLSSSPNI NSYSESGVYSQESLTSSLSTTPQGKRIMSDIFPTFG SKPCPTRLSSAKKKISHIAEQSPSAGSSSNPQQISS FDFTTTKALSEDSVVVVGKGVFGSLSSAPATCSQ SVISSVENGDTFSIKQSIEPPSGIYGRSVQQNISSYL DVENEKDAKVSISKSTYNKMRQKRKEEKELFHN KDCEKKEKNSWERMRHTGTEKMASESETPTGAI SQYKERMPSVTHSPEIMDLSELRPFSKPEIALTEA LRLLADEDWEKKIEGLNFIRCLAAFHSEILNTKL HETNFAVVQEVKNLRSGVSRAAVVCLSDLFTYL KKSMDQELDTTVKVLLHKAGESNTFIREDVDKA LRAMVNNVTPARAVVSLINGGQRYYGRKMLFF MMCHPNFEKMLEKYVPSKDLPYIKDSVRNLQQK GLGEIPLDTPSAKGRRSHTGSVGNTRSSSVSRDA FNSAERAVTEVREVTRKSVPRNSLESAEYLKLIT GLLNAKDFRDRINGIKQLLSDTENNQDLVVGNIV KIFDAFKSRLHDSNSKVNLVALETMHKMIPLLRD HLSPIINMLIPAIVDNNLNSKNPGIYAAATNVVQA LSQHVDNYLLLQPFCTKAQFLNGKAKQDMTEKL ADIVTELYQRKPHATEQKVLVVLWHLLGNMTN SGSLPGAGGNIRTATAKLSKALFAQMGQNLLNQ AASQPPHIKKSLEELLDMTILNEL
3517	A	1449	252	QDLKPVLDREYLAIYLKMVFFTCNACGESVKKI QVEKHVSVCRNCECLSCIDCGKDFWGDDYKNH VKCISEDQKYGGKGY/EKVKTHKGD/ASKQQAW IQKISELIK\RPNVSPKVRELLEQISAFDNVPQ\KK AKFQNWMKNSLKVHNESILDQVWNIFSEASNSE PVNKEQDQRPLHPVANPHAEISTKVPASKVKDA VEQQGEVKKNKRERKEERQKKRKREKKELKLE NHQENSRNQKPKKRKKGQEADLEAGGEEVPEA NGSAGKRSKKKKQRKDSASEEEARVGAGKRKR RHSKVETDSKKKKMKLPEHPEGGEPEDDEAPAK GKFNWKGTIKAILKQAPDNEITIKKLRKKVLAQY YTVTDEHHRSEEELLVIFNKKISKNPTFKLLKDK VKLVK
3518	A	3	635	APDSNARNDHFDACSLRVQAGLSSAGPALGNSG LAALMASPSKAVIVPGNGGGDVTTHGWYGWVK KELEKIPGFQCLAKNMPDPITARESIWLPFMETEL HCDEKTIIIGHSSGAIAAMRYAETHRVYAIVLVSA YTSDLGDENERASGYFTRPWQWEKIKANCPYIV QFGSTDDPFLPWKEQQEVAD\SWKPNCTNSLTV ATFRTQSFMN
3519	A	81	2277	VRETRREMAMAMSDSGASRLRRQLESGGFEARL YVKQLSQQSDGDRDLQEHRQRIQALAEETAQNL KRNVYQNYRQFIETAREISYLESEMYQLSHLLTE QKSSLESIPLTLLPAAAAAGAAAASGGEEGVGGA GGRDHLRGQAGFFSTPGGASRDGSGPGEEGKQR TLTTLLEKVEGCRHLLETPGQYLVYNGDLVEYD ADHMAQLQRVHGFLMNDCLLVATWLPQRRGM

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SEQ ID NO:	Method	Predicted beginning nucleotide location corresponding to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion
				YRYNALYSLDGLAVVNVKDNPPMKDMFKLLMF PENRIFQAENAKIKREWLEVLEDTKRALSEKRRR EQEEAAAPRGPPQVTSKATNPFEDDEEEEPAVPE VEEEKVDLSMEWIQELPEDLDVCIAQRDFEGAV DLLDKLNHYLEDKPSPPPVKELRAKVEERVRQL TEVLVFELSPDRSLRGGPKATRRAVSQLIRLGQC TKACELFLRNRAAAVHTAIRQLRIEGATLLYIHK LCHVFFTSLLETAREFEIDFAGTDSGCYSAFVVW ARSAMGMFVDAFSKQVFDSKESLSTAAECVKVA KEHCQQLGDIGLDLTFIIHALLVKDIQGALHSYK EIIEATKHRNSEEMWRRMNLMTPEALGKLKEE MKSCGVSNFEQYTGDDCWVNLSYTVVAFTKQT MGFLEEALKLYFPELHMVLLESLVEIILVAVQHV DYSLRCEQDPEKKAFIRQNASFLYETVL\PVVEK RFEEGVGKPAKQLQDLRNASRLIRVNPESTTSVV
3520	A	1706	540	FVAHLAWPWRADGDMEDGVLNEGFLVKRGHIV HNWKARWFILRQNTLVYYKLEGGRRVTPPKGRI LLDGCTITCPCLEYENRPLLIKLKTQTSTEYFLEA CSREE/RRDAWAFE\ITGAIHAGQARGKVQQLHS LRNSFKLPPHISLHRIVDKMHDSNTGIRSSPNMEQ GSTYKKTFLGSSLVDWLISNSFTASRLEAVTLAS MLMEENFLRPVGVRSMGAIRSGDLAEQFLDDST ALYTFAESYKKKISPKEEISLSTVELSGTVVKQGY LAKQGHKRKNWKVRRFVLRKDPAFLHYYDPSK EENRPVGGFSLRGSLVSALEDNGVPTGVKGNVQ GNLFKVITK\DDTHYYIQA\SSKAE\RAE\WIGSLS KSLNMNKDPEGTPDSLPSLPR
3521	A	3	3063	HASVSLSLGCPRPCADTPGPQPQPMDLRVGQRPP VEPPPEPTLLALQRPQRLHHHLFLAGLQQQRSVE PMRVKMELPACGATLSLVPSLPAFSIPRHQSQSST PCPFLGCRPCPQLSMDTPMPELQEAPQEQELRQL LHKDKSKRSAVASSVVKQKLAEVILKKQQAALE RTVHPNSPGIPYRTLEPLETEGATRSMLSSFLPPV PSLPSDPPEHFPLRKTVSEPNLKLRYKPKKSLERR KNPLLRKESAPPSLRRRPAETLGDSSPSSSSTPAS GCSSPNDSEHGPNPILGSEALLGQRLRLQETSVAP FALPTVSLLPAITLGLPAPARADSDRRTHPTLGPR GPILGSPHTPLFLPHGLEPEAGGTLPSRLQPILLLD PSGSHAPLLTVPGLGPLPFHFAQSLMTTERLSGSG LHWPLSRTRSEPLPPSATAPPPPGPMQPRLEQLKT HVQVIKRSAKPSEKPRLRQIPSAEDLETDGGGPG QVVDDGLEHRELGHGQPEARGPAPLQQHPQVLL WEQQRLAGRLPRGSTGDTVLLPLAQGGHRPLSR AQSSPAAPASLSAPEPASQARVLSSSETPARTLPF TTGLIYDSVMLKHQCSCGDNSRHPEHAGRIQSIW SRLQERGLRSQCECLRGRKASLEELQSVHSERHV LLYGTNPLSRLKLDNGKLAGLLAQRMFVMLPCG GVGVDTDTIWNELHSSNAARWAAGSVTDLAFK VASRELKNGFAVVRPPGHHADHSTAMGFCFFNS VAIACRQLQQSKASKILIVDWDVHHGNGTQQT FYQDPSVLYISLHRHDDGNFFPGSGAVDEVGAGS GEGFNVNVAWAGGLDPPMGDPEYLAAFRIVVM PIAREFSPDLVLVSAGFDAAEGHPAPLGGYHVSA KCFGYMTQQLMNLAGGAVVLALEGGHDLTAIC DASEACVAALLGNRVDPLSEEGWKQKPNLNAIR

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SEQ ID	Method	Predicted	Predicted end	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine,
NO:		beginning nucleotide	nucleotide {	E=Glutamic Acid, F=rhenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine,
		location	corresponding	N=Asparagine, P=Proline, O=Glutamine, R=Arginine, S=Serine,
		corresponding	to last amino	T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine,
		to first amino acid residue of	acid residue of peptide	X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion
		peptide sequence	sequence	
				SLEA\VIRVHSKYWGCMQRLASCPDSWVPRVPG ADKEEVEAVTALASLSVGILAEDRPSEQLVEEEE
				PMNL
3522	Α	9	602	KMAALGEPVRLERDICRAIELLEKLQRSGEVPPQ
ĺ				KLQALQRVLQSEFCNAVREVYEHVYETVDISSSP
				EVRANATAKATVAAFAASEGHSHPRVVELPKTE EGLGFNIMGGKEQNSPIYISRIIP/GGIADRHGGLK
				RGDQLLSVNGVSVEGEHHEKAVELLKAAQGKV
			İ	KLVVRYTPKVLEEMESRFEKMRSAKRRQQT
3523	A	645	1465	IMAETSLLEAGASAASTAAALENLQVEASCSVCL
				EYLKEPVIECGHNFCKACITRWWEDLERDFPCP
				VCRKTSRYRSLRPNRQLGSMVEIAKQL\RPSSGRS
		1		GMRASAPQHHEALSLFCYEDQEAVCLICAISHTH
				RAHTVVPLDDATQEYKEKLQKCLEA\LNQKLQEI
	İ			TRCKSSEEKKPGELKRLVESRRQQILREFEELHRR
	1			LDEEQQVLLSRLEEEEQDILQRLRENAAHLGDKR RDLAHLAAEVEGKCLQSGFEMLKVRPLPLHSPS
	1		ļ	G
3524	A	3	698	PMVRHEAGEALGAIGDPEVLEILKQYSSDPVIEV
***	1			AETCQLAVRRLEWLQQHGGEPAAGPYLSVDPAP
				PAEER\DVGRLREALLDESRPLFERYRAMFALRN
				AGGEEAALALAEGLHCGSALFRHEVGYVLGQLQ
1	-	1		HEAAVPQLAAALARCTENPMVRHECAEALGAIA
		1		RPACLAALQAHADDPERVVRE\SCKVALDMYEH
2525	<u> </u>	1452	694	ETGRAFQYADGLEQLRGAPSLGPNPHPELPEDS EGLQRPEYLVASAAGFQGLAWGGEGRGRAGCS
3525	A	1432	094	SSGFRDAEPLLLSCPGRNEPLKKERLKWKSDYP
				MTDGQLRSKRDEFWDTAPAFEGRKEIWDALKA
	-			AAYAAEANDHELAQAILDGASITLPHGTLCECY
				DELGNRYQLPIYCLSPPVNLLLEHTEEESLEPPEP
ł		ł	1	PPSVRREFPLKVRLSTGKDVRLSASLPDTVGQLK
				RQLHAQE/GTPKPSWQRWFFSGKLLTDRTRLQET
				KIQKDFVIQVIINQPPPPQD
3526	A	123	3441	PGNEGLGLAADHNEDLGHLSADAPWPAVTMAP
}			ļ	RKRSHHGLGFLCCFGGSDIPEINLRDNHPLQFME FSSPIPNAEELNIRFAELVDELDLTDKNREAMFAL
			1	PPEKKWQIYCSKKKEQEDPNKLATSWPDYYIDRI
	1			NSMAAMQSLYAFDEEETEMRNQVVEDLKTALR
				TOPMRFVTRFIELEGLTCLLNFLRSMDHATCESRI
İ				HTSLIGCIIALMNNSQGRAHVLAQPEAISTIAQSL
ĺ			-	RTENSKTKVAVLEILGAVCLVPGGHKKVLQAML
	1			HYQVYAAERTRFQTLLNELDRSLGRYRDEVNLK
	1			TAIMSFINAVLNAGAGEDNLEFRLHLRYEFLMLG
				IQPVIDKLRQHENAILDKHLDFFEMVRNEDDLEL
1	1		1	ARRFDMVHIDTKSASQMFELIHKKLKYTEAYPC LLSVLHHCLQMPYKRNGGYFQQWQLLDRILQQI
			1	LLSVLHHCLQMPYKKNGGYFQQWQLLDKILQQI VLQDERGVDPDLAPLENFNVKNIVNMLINENEV
				KQWRDQAEKFRKEHMELVSRLERKERECETKTL
			1	EKEEMMRT\LNKMKDKLARESQELRQARGQVA
				ELVAQLSELSTGPVSSPPPPGGPLTLSSSMTTNDL
				PPPPPPLPFACCPPPPPPPPLPPGGPPTPPGAPPCLG
				MGLPLPQDPYPSSDVPLRKKRVPQPSHPLKSFNW
				VKLNEERVPGTVWNEIDDMQVFRILDLEDFEKM
				FSAYQRHQELITNPSQQKELGSTEDIYLASRKVK
L	l			ELSVIDGRRAQNCIILLSKLKLSNEEIRQAILKMD

SEQ ID NO:	Method	Predicted beginning nucleotide location corresponding to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide 'sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion
				EQEDLAKDMLEQLLKFIPEKSDIDLLEEHKHEIER MARADRFLYEMSRIDHYQQRLQALFFKKKFQER LAEAKPKVEAILLASRELVRSKRLRQMLEVILAI GNFMNKGQRGGAYGFRVASLNKIADTKSSIDRN ISLLHYLIMILEKHFPDILNMPSELQHLPEAAKVN LAELEKEVGNLRRGLRAVEVELEYQRRQVREPS DKFVPVMSDFITVSSFSFSELEDQLNEARDKFAK ALMHFGEHDSKMQPDEFFGIFDTFLQAFSEARQD LEAMRRKEEEERRARMEAMLKEQRERERWQR QRKVLAAGSSLEEGGEFDDLVSALRSGEVFDKD LCKLKRSRKRSGSQALEVTRERAINRLNY
3527	A	1445	714	LLGTRMLAGQLEARDPKEGTHPEDPCPGAGAV MEKTAVAAEVLTEDCNTGEMPPLQQQIIRLHQE LGRQKSLWADVHGKLRSHIDALREQNMELREKL RALQLQRWKARKKSAASPHAGQESHTLALEPAF GKISPLSADEETIPKYAGHKN\QSGHSSWGQRSSS NNSAPPKPMSLKIERISSWKTPPQENRDKNLSRR RQDRRATPTGRPTPCAERRG\VSEDGKVASDTCV TLHWPLGKFRFR
3528	A	484	1777	RISKIQVYYSTGYSSRKMNPTLGLAIFLAVLLTVK GLLKPSFSPRNYKALSEVQGWKQRMAAKELAR QNMDLGFKLLKKLAFYNPGRNIFLSPLSISTAFS MLCLGAQDSTLDEIKQGFNFRKMPEKDLHEGFH YIIHELTQKTQDLKLSIGNTLFIDQRLQPQRKFLE DAKNFYSAETILTNFQNLEMAQKQINDFI/ESKTH GKINNLIENIDPGTVMLLANYIFFRARWKHEFDP NVTKEEDFFLEKNSSVKVPMMFRSGIYQVGYDD KLSCTILEIPYQKNITAIFILPDEGKLKHLEKGLQV DTFSRWKTLLSRRVVDVSVPRLHMTGTFDLKKT LSYIGVSKIFEEHGDLTKIAPHRSLKVGEAVNKA ELKMDERGTEGAAGTGAQTLPMETPLVVKIDKP YLLLIYSEKIPSVLFLGKIVNPIGK
3529	A		5684	VSSVSHENPTEVFEDGENPPSSRSSESGFTEFIQY QADRTDDIDRELSEGQGAAAIPIGSTSSETETAST VGSEETIIQTPSVVTQGTATRSRKTAQKTAMQCC LEYVQQFLTRLINLYIIQNNSFSQSLATEHQGDLG REQGETSKWDRNSQGDVKEKNISKQKTSKEYLS AFLAACQLFLECSSFPVYIAEGNHTSELRSEKLET DCEHVQPPQWLQTLMNACSQASDFSVQSVAISL VMDLVGLTQSVAMVTGENINSVEPAQPLSPNQG RVAVVIRPPLTQGNLRYIAEKTEFFKHVALTLWD QLGDGTPQHHQKSVELFYQLHNLVPSSSICEDVI SQQLTHKDKKIRMEAHAKFAVLWHLTRDLHINK SSSFVRSFDRSLFIMLDSLNSLDGSTSSVGQAWL NQVLQRHDIARVLEPLLLLLLHPKTQRVSVQRV QAERYWNKSPCYPGEESDKHFMQNFACSNVSQ VQLITSKGNGEKPLTMDEIENFSLTVNPLSDRLSL LSTSSETIPMVVSDFDLPDQQIEILQSSDSGCSQSS AGDNLSYEVDPETVNAQEDSQMPKESSPDDDVQ QVVFDLICKVVSGLEVESASVTSQLEIEAMPPKC SDIDPDEETIKIEDDSIQQSQNALLSNESSQFLSVS AEGGHECVANGISRNSSSPCISGTTHTLHDSSVAS IETKSRQRSHSSIQFSFKEKLSEKVSEKETIVKESG KQPGAKPKVKLARKKDDDKKKSSNEKLKQTSV FFSDGLDLENWYSCGEGDISEIESDMGSPGSRKSP

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000	1.54	I 15-141-14-14	Danding 3	Amino acid sequence (A≈Alanine C=Cysteine, D=Aspartic Acid,
SEQ ID NO:	Method	Predicted	Predicted end nucleotide	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine,
NO:		beginning nucleotide	location	I=Isoleucine, K=Lysine, L=Leucine, M=Methionine,
	ł	location	corresponding	N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine,
	1	corresponding	to last amino	T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine,
		to first amino	acid residue of	X=Unknown, *=Stop codon, /=possible nucleotide deletion,
		acid residue of	peptide	\=possible nucleotide insertion
		peptide sequence	sequence	
	1			NFNIHPLYQHVLLYLQLYDSSRTLYAFSAIKAILK
				TNPIAFVNAISTTSVNNAYTPQLSLLQNLLARHRI
				SVMGKDFYSHIPVDSNHNFRSSMYIEILISLCLYY
				MRSHYPTHVKVTAQDLIGNRNMQMMSIEILTLL
				FTELAKVIESSAKGFPSFISDMLSKCKVQKVILHC
			1	LLSSIFSAQKWHSEKMAGKNLVAVEEGFSEDSLI
				NFSEDEFDNGSTLQSQLLKVLQRLIV\LEHRVM\T
			İ	IPEE\NETGFDFVVS\DLEHISPHQPMTSLQYLHAQ
			1	SITCQGMFLCAVIRA\LHQHCACKMHPQWIGLIT
				STLPYMGKVLQRVVVSVTLQLCRNLDNLIQQYK
	1		İ	YETGLSDSRPLWMASIIPPDMILTLLEGITAIIHYC
				LLDPTTQYHQLLVSVDQKHLFEARSGILSILHMI
				MSSVTLLWSILHQADSSEKMTIAASASLTTINLG
			İ	ATKNLRQQILELLGPISMNHGVHFMAAIAFVWN
				ERRONKTTTRTKVIPAASEEQLLLVELVRSISVM
			ļ	RAETVIQTVKEVLKQPPAIAKDKKHLSLEVCML
				QFFYAYIQRIPVPNLVDSWASLLILLKDSIQLSLP
			[APGQFLILGVLNEFIMKNPSLENKKDQRDLQDVT
	ŀ			HKIVDAIGAIAGSSLEQTTWLRRNLEVKPSPKIM
			}	VDGTNLESDVEDMLSPAMETANITPSVYSVHAL
ļ	1			TLLSEVLAHLLDMVFYSDEKERVIPLLVNIMHYV
·				VPYLRNHSAHNAPSYRACVQLLSSLSGYQYTRR
				AWKKEAFDLFMDPSFFQMDASCVNHWRAIMDN
	Í		1	LMTHDKTTFRDLMTRVAVAQSSSLNLFANRDVE
1				LEQRAMLLKRLAFAIFSSEIDQYQKYLPDIQERLV
l	ì	1		ESLRLPQVPTLHSQVFLFFRVLLLRMSPQHLTSL
			<u> </u>	WPTMITELVQVFLLMEQELTADEDISRTSGPSVA
				GLETTYTGGNGFSTSYNSQRWLNLYLSACKFLD
				LALALPSENLPQFQMYRWAFIPEASDDSGLEVRR
ĺ	[QGIHQREFKPYVVRLAKLLRKRAKKNPEEDNSG
				RTLGWEPGHLLLTICTVRSMEQLLPFFNVLSQVF
 	}		}	NSKVTSRCGGHSGSPILYSNAFPNKDMKLENHKP
-		ļ		CSSKARQKIEEMVEKDFLEGMIKT
3530	A	1	5684	VSSVSHENPTEVFEDGENPPSSRSSESGFTEFIQY
2230	^	1	3004	QADRTDDIDRELSEGQGAAAIPIGSTSSETETAST
[[VGSEETIIQTPSVVTQGTATRSRKTAQKTAMQCC
ļ	1			LEYVQQFLTRLINLYIIQNNSFSQSLATEHQGDLG
f				REQGETSKWDRNSQGDVKEKNISKQKTSKEYLS
				AFLAACQLFLECSSFPVYIAEGNHTSELRSEKLET
				DCEHVQPPQWLQTLMNACSQASDFSVQSVAISL
		ļ		VMDLVGLTQSVAMVTGENINSVEPAQPLSPNQG
ĺ	ļ			RVAVVIRPPLTQGNLRYIAEKTEFFKHVALTLWD
				QLGDGTPQHHQKSVELFYQLHNLVPSSSICEDVI
		1		SQQLTHKDKKIRMEAHAKFAVLWHLTRDLHINK
ļ	}		}	SSSFVRSFDRSLFIMLDSLNSLDGSTSSVGQAWL
	-			NQVLQRHDIARVLEPLLLLLHPKTQRVSVQRV
	1			
[1			QAERYWNKSPCYPGEESDKHFMQNFACSNVSQ
	[VQLITSKGNGEKPLTMDEIENFSLTVNPLSDRLSL
1				LSTSSETIPMVVSDFDLPDQQIEILQSSDSGCSQSS
]	}			AGDNLSYEVDPETVNAQEDSQMPKESSPDDDVQ
}				QVVFDLICKVVSGLEVESASVTSQLEIEAMPPKC
	}	'		SDIDPDEETIKIEDDSIQQSQNALLSNESSQFLSVS
	1		1	AEGGHECVANGISRNSSSPCISGTTHTLHDSSVAS
				IETKSRQRSHSSIQFSFKEKLSEKVSEKETIVKESG
L			<u> </u>	KQPGAKPKVKLARKKDDDKKKSSNEKLKQTSV
		J		

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SEQ ID NO:	Method	Predicted beginning nucleotide location corresponding to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion
				FFSDGLDLENWYSCGEGDISEIESDMGSPGSRKSP NFNIHPLYQHVLLYLQLYDSSRTLYAFSAIKAILK TNPIAFVNAISTTSVNNAYTPQLSLLQNLLARHRI SVMGKDFYSHIPVDSNHNFRSSMYIEILISLCLYY MRSHYPTHVKVTAQDLIGNRNMQMMSIEILTLL FTELAKVIESSAKGFPSFISDMLSKCKVQKVILHC LLSSIFSAQKWHSEKMAGKNLVAVEEGFSEDSLI NFSEDEFDNGSTLQSQLLKVLQRLIVLEHRVM\T IPEE\NETGFDFVVS\DLEHISPHQPMTSLQYLHAQ SITCQGMFLCAVIRA\LHQHCACKMHPQWIGLIT STLPYMGKVLQRVVVSVTLQLCRNLDNLIQQYK YETGLSDSRPLWMASIIPPDMILTLLEGITAIIHYC LLDPTTQYHQLLVSVDQKHLFEARSGILSILHMI MSSVTLLWSILHQADSSEKMTIAASASLTTINLG ATKNLRQQILELLGPISMNHGVHFMAAIAFVWN ERRQNKTTTRTKVIPAASEEQLLLVELVRSISVM RAETVIQTVKEVLKQPPAIAKDKKHLSLEVCML QFFYAYIQRIPVPNLVDSWASLLILLKDSIQLSLP APGQFLILGVLNEFIMKNPSLENKKDQRDLQDVT HKIVDAIGAIAGSSLEQTTWLRRNLEVKPSPKIM VDGTNLESDVEDMLSPAMETANITPSVYSVHAL TLLSEVLAHLLDMVFYSDEKERVIPLLVNIMHYV VPYLRNHSAHNAPSYRACVQLLSSLSGYQYTRR AWKKEAFDLFMDPSFFQMDASCVNHWRAIMDN LMTHDKTTFRDLMTRVAVAQSSSLNLFANRDVE LEQRAMLLKRLAFAIFSSEIDQYQKYLPDIQERLV ESLRLPQVPTLHSQVFLFFRVLLRMSPQHLTSL WPTMITELVQVFLLMEQELTADEDISRTSGPSVA GLETTYTGGNGFSTSYNSQRWLNLYLSACKFLD LALALPSENLPQFQMYRWAFIPEASDDSGLEVRR QGIHQREFKPYVVRLAKLLRKRAKKNPEEDNSG RTLGWEPGHLLLTICTVRSMEQLLPFFNVLSQVF NSKVTSRCGGHSGSPILYSNAFPNKDMKLENHKP CSSKARQKIEEMVEKDFLEGMIKT
3531	A	553	2470	LISPSPALSSQDPALSLKENLEDISGWGLPEARSK ESVSFKDVAVDFTQEEWGQLDSPQRALYRDVM LENYQNLLALGPPLHKPDVISHLERGEEPWSMQ REVPRGPCPEWELKAVPSQQQGICKEEPAQEPIM ERPLGGAQAWGRQAGALQRSQAAP\GR\RTCHG LGRP\VEEFPLRCPLFAQQRVPEGGPLLDTRKNV QATEGRTKAPARLCAGENASTPSEPEKFPQVRRQ RGAGAGEGEFVCGECGKAFRQSSSLTLHRRWHS REKAYKCDECGKAFTWSTNLLEHRRIHTGEKPFF CGECGKAFSCHSSLNVHQRIHTGERPYKCSACEK AFSCSSLLSMHLRVHTGEKPYRCGECGKAFNQR THLTRHHRIHTGEKPYQCGSCGKAFTCHSSLTVH EKIHSGDKPFKCSDCEKAFNSRSRLTLHQRTHTG EKPFKCADCGKGFSCHAYLLVHRRIHSGEKPFKC NECGKAFSSHAYLIVHRRIHTGEKPFDCSQCWKA FSCHSSLIVHQRIHTGEKPYKCSECGRAFSQNHCL IKHQKIHSGEKSFKCEKCGEMFNWSSHLTEHQRL HSEGKPLAIQFNKHLLSTYYVPGSLLGAGDAGLR DVDPIDALDVAKLLCVVPPRAGRNFSLGSKPRN
3532	A	3931	317	HRELQDSPSAEPPAGSMPLRHWGMARGSKPVGD GAQPMAAMGGLKVLLHWAGPGGGEPWVTFSES

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SEQ ID NO:	Method	Predicted beginning nucleotide location corresponding to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion
				SLTAEEVCIHIAHKVGITPPCFNLFALFDAQAQV WLPPNHILEIPRDASLMLYFVRHRFYSR\NWHGM NPREPAVYRCGPPGTEASSDQTAQGMQLLDPAS FEYLFEQGKHEFVNDVASLWELSTEEEIHHFKNE SLGMAFLHLCHLALRHGIPLEEVAKKTSFKDCIP RSFRRHIRQHSALTRLRLRNVFRRFLRDFQPGRLS QQMVMVKYLATLERLAPRFGTERVPVCHLRLLA QAEGEPCYIRDSGVAPTDPGPESAAGPPTHEVLV TGTGGIQWWPVEEEVNKEEGSSGSSGRNPQASL FGKKAKAHKAFGQPADRPREPLGAYFCDFRDIT HVGLKEHCVSIHRQDNKCLELSLPSRAAALSFVS LVDGYFRLTADSSHYLCHEVAPPRLVMSIRDGIH GPLLEPFVQAKLRPEDGLYLIHWSTSHPYRLILTV AQRSQAPDGMQSLRLRKFPIEQQDGAFVLEGWG RSFPSVRELGAALQGCLLRAGDDCFSLRRCCLPQ PGETSNLIIMRGARASPRTLNLSQLSFHRVDQKEI TQLSHLGQGTRTNVYEGRLRVEGSGDPEEGKMD DEDPLVPGRDRGQELRVVLKVLDPSHHDIALAF YETASLMSQVSHTHLAFVHGVCVRGPENIMVTE YVEHGPLDVWLRRERGHVPMAWKMVVAQQLA SALSYLENKNLVHGNVCGRNILLARLGLAEGTSP FIKLSDPGVGLGALSREERVERIPWLAPECLPGG ANSLSTAMDKWGFGATLLEICFDGEAPLQSRSPS EKEHFYQRQHRLPEPSCPQLATLTSQCLTYEPTQ RPSFRTILRDLTRLQPHNLADVLTVNPDSPASDPT VFHKRYLKKIRDLGEGHFGKVSLYCYDPTNDGT GEMVAVKALKADCGPQHRSGWKQEIDILRTLYH EHIIKYKGCCEDQGEKSLQLVMEYVPLGSLRDYL PRHSIGLAQLLLFAQQICEGMAYLHAQHYIHRDL AARNVLLDNDRLVKIGDFGLAKAVPEGHEYYRV REDGDSPVFWYAPECLKEYKFYYASDVWSFGVT LYELLTHCDSSQSPPTKFLELIGIAQGQMTVLRLT ELLERGERLPRPDKCPCEVYHLMKNCWETEASF RPTFENLIPILKTVHEKYQGQAPSVFSVC
3533	A	182	3465	FRWLDFFRGSINSQFEFGRKKENMTSPAKFKKDK EIIAEYDTQVKEIRAQLTEQMKCLDQQCELRVQL LQDLQDFFRKKAEIEMDYSRNLEKLAERFLAKT RSTKDQQFKKDQNVLSPVNCWNLLLNQVKRES RDHTTLSDIYLNNIIPRFVQVSEDSGRLFKKSKEV GQQLQDDLMKVLNELYSVMKTYHMYNADSISA QSKLKEAEKQEEKQIGKSVKQEDRQTPRSPDSTA NVRIEEKHVRSSVKKIEKMKEKRQAKYTENKL KAIKARNEYLLALEATNASVFKYYIHDLSDLIDQ CCDLGYHASLNRALRTFLSAELNLEQSKHEGLD AIENAVENLDATSDKQRLMEMYNNVFCPPMKFE FQPHMGDMASQLCAQQPVQSELLQRCLQLQSRL STLKIENEEVKKTMEATLQTIQDIVTVEDFDVSD CFQYSNSMESVKSTVSETFMSKPSIAKRANQQE TEQFYFTKMKEYLEGRNLITKLQAKHDLLQKTL GESQRTDCSLARRSSTVRKQDSSQAIPLVVESCIR FISRHGLQHEGIFRVSGSQVEVNDIKNAFERGEDP LAGDQNDHDMDSIAGVLKLYFRGLEHPLFPKDIF HDLMACVTMDNLQERALHIRKVLLVLPKTTLII MRYLFAFLNHLSQFSEENMMDPYNLAICFGPSL MSVPEGHDQVSCQAHVNELIKTIIIQHENIFPSPRE

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SEQ ID NO:	Method	Predicted beginning nucleotide location corresponding to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion
				LEGPVYSRGGSMEDYCDSPHGETTSVEDSTQDV TAEHHTSDDECEPIEAIAKFDYVGRTARELSFKK GASLLLYQRASDDWWEGRHNGIDGLIPHQYIVV QDTEDGVVERSSPKSEIEVISEPPEEKVTARAGAS CPSGGHVADIYLANINKQRKRPESGSIRKTFRSDS HGLSSSLTDSSSPGVGASCRPSSQPIMSQSLPKEG PDKCSISGHGSLNSISRHSSLKNRLDSPQIRKTAT AGRSKSFDNHRPMDPEVIAQDIEATMNSALNELR ELERQSSVKHTPDVVLDTLEPLKTSPVVAPTSEPS SPLHTQLLKDPEPAFQRSASTAGDIACAFRPVKS VKMAAPVKPPAT\RPKPT\VFPKTNATSPGVNSST
3534	A	:	2640	FRFVCPASRRPAAGLRDAASSAPRGMASEGPRE PESEGIKLSADVKPFVPRFAGLNVAWLESSEACV FPSSAATYYPFVQEPPVTEQKIYTEDMAFGASTFP PQYLSSEITLHPYAYSPYTLDSTQNVYSVPGSQY LYNQPSCYRGFQTVKHRNENTCPLPQEMKALFK KKTYDEKKTYDQQKFDSERADGTISSEIKSARGS HHLSIYAENSLKSDGYHKRTDRKSRIIAKNVSTS KPEFEFTTLDFPELQGAENNMSEIQKQPKWGPVH SVSTDISLLREVVKPAAVLSKGEIVVKNNPNESV TANAATNSPSCTRELSWTPMGYVVRQTLSTELS AAPKNVTSMINLKTIASSADPKNVSIPSSEALSSD PSYNKEKHIIHPTQKSKASQGSDLEQNEASRKNK KKKEKSTSKYEVLTVQEPPRIEDAEEFPNLAVAS ERRDRIETPKFQSKQQPQDNFKNNVKKSQLPVQL DLGGMLTALEKKQHSQHAKQSSKPVVVSVGAV PVLSKECASGERGRRMSQMKTPHNPLDSSAPLM KKGKQREIPKAKKPTSLKKIILKERQERKQRLQE NAVSPAFTSDDTQDGESGGDDQFPEQAELSGPEG MDELISTPSVEDKSEEPPGTELQRDTEASHLAPN HTTFPKIHSRRFRDYCSQMLSKEVDACVTDLLKE LVRFQDRMYQKDPVKAKTKRRLVLGLREVLKH LKLKKLKCVIISPNCEKIQSKGGLDDTLHTIIDYA
3535	A	1747	983	CEQNIPFVFALNRKALGRSLNKAVPVSVVGIFSY DGAQDQFHKMVELTVAARQAYKTMLENVQQE LVGEP\SLRHLPAYPHRAPAALQKMAPQP/VKEK EEPHYIEIWKKHLEAYSGCTLELEESLEASTSQM MNLNL LFQFQVCRSVLSPRAAGCTWSLAPRSRGAAGSPR RYRGPQPQPAPPSALPNSRPSPVASGREMVVLSV PAEVTVILLDIEGTTTPIAFVKDILFPYIEENVKEY LQTHWEEEECQQDVSLLRKQV\FADVVPAVRKW
3536	A	3	1302	REAGMKVYIYSSGSVEAQKLLFGHSTEGDILELV DGHFDTKIGHKVESESYRKIADSIGCSTNNILFLT DVTREASAAEEADVHVAVVVRPGNAGLTDDEK TYYSLITSFSELYLPSST GRPPTAPHTGRPPTANRGDPRLDLKRGCARLLTS IESRGRPAASAGLRRDRCALRRWPLRRAPLARAT RRRAGSPRRCAPRPRACPQGWSRARHQPGGLCL LLLLLCQFMEDRSAQAGNCWLRQAKNGRCQVL YKTELSKEECCSTGRLSTSWTEEDVNDNTLFKW MIFNGGAPNCIPCKETCENVDCGPGKKCRMNKK
				NKPRCVCAPDCSNITWKGPVCGLDGKTYRNECA LLKARCKEQPELEVQYQGRCKKTCRDVFCPGSS

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				TCV\VDQTNNAYCVTCNRICPEPASSEQYLCGND GVTYS\SACHLRKATCLLGRSIGLAYEGKCIKAK SCEDIQCTGGKKCLWDFKVGRGRCSLCDELCPD SKSDEPVCASDNATYASECAMKEAACSSGVLLE VKHSGSCNSISEDTEEEEEDEDQDYSFPISSILEW
3537	A	285	2123	IGLFLQVAPLSVMAKSCPSVCRCDAGFIYCNDRF LTSIPTGIPEDATTLYLQNNQINNAGIPSDLKNLL KVERIYLYHNSLDEFPTNLPKYVKELHLQENNIR TITYDSLSKIPYLEELHLDDNSVSAVSIEEGAFRD SNYLRLLFLSRNHLSTIPWGLPRTIEELRLDDNRIS TISSPSLQGLTSLKRLVLDGNLLNNHGLGDKVFF NLVNLTELSLVRNSLTAAPVNLPGTNLRKLYLQ DNHINRVPPNAFSYLRQLYRLDMSNNNLSNLPQ GIFDDLDNITQLILRNNPWYCGCKMKWVRDWL QSLPVKVNVRGLMCQAPEKVRGMAIKDLNAELF DCKDSGIVSTIQITTAIPNTVYPAQGQWPAPVTK QPDIKNPKLTKDHQTTGSPSRKTITITVKSVTSDTI HISWKLALPMTALRLSWLKLGHSPAFGSITETIVT GERSEYLVTALEPDSPYKVCMVPMETSNLYLFD ETPVCIETETAPLRMYNPTTTLNREQEKEPYKNP NLPLAAIIGGAVALVTIALLALVCWYVHRNGSLF SRNCAYSKGRRRKDDYAEAGTKKDNSILEIRETS FQMLPISNEPISKEEFVIHTIFPPNGMNLYKNNH
3538	A	877	6184	WNVKPSLLVVQLFKFSDKEEHEQNDSISGKTGET GVEEMIATRKVEQDSKETVKLSHEDDHILEDAGS SDISSDAACTNPNKTENSLVGLPSCVDEVTECNL ELKDTMGIADKTENTLERNKIEPLGYCEDAESNR QLESTEFNKSNLEVVDTSTFGPESNILENAICDVP DQNSKQLNAIESTKIESHETANLQDDRNSQSSSV SYLESKSVKSKHTKPVIHSKQNMTTDAPKKIVAA KYEVIHSKTKVNVKSVKRNTDVPESQQNFHRPV KVRKKQIDKEPKIQSCNSGVKSVKNQAHSVLKK TLQDQTLVQIFKPLTHSLSDKSHAHPGCLKEPHH PAQTGHVSHSSQKQCHKPQQQAPAMKTNSHVK EELEHPGVEHFKEEDKLKLKKPEKNLQPRQRRSS KSFSLDEPPLFIPDNIATIRREGSDHSSSFESKYMW TPSKQCGFCKKPHGNRFMVGCGRCDDWFHGDC VGLSLSQAQQMGEEDKEYVCVKCCAEEDKKTEI LDPDTLENQATVEFHSGDKTMECEKLGLSKHTT NDRTKYIDDTVKHKVKILKRESGEGRNSSDCRD NEIKKWQLAPLRKMGQPVLPRRSSEEKSEKIPKE STTVTCTGEKASKPGTHEKQEMKKKKVEKGVL NVHPAASASKPSADQIRQSVRHSLKDILMKRLTD SNLKVPEEKAAKVATKIEKELFSFFRDTDAKYKN KYRSLMFNLKDPKNNILFKKVLKGEVTPDHLIR MSPEELASKELAAWRRENRHTIEMIEKEQREVE RRPITKITHKGEIEIESDAPMKEQEAAMEIQEPAA NKSLEKPEGSEK\RKEEVDSMSKDTTSQHRQHLF DLNCKICIGRMAPPVDDLSPKKVKVVVGVARKH SDNEAESIADALSSTSNILASEFFEEEKQESPKSTF SPAPRPEMPGTVEVESTFLARLNFIWKGFINMPS VAKFVTKAYPVSGSPEYLTEDLPDSIQVGGRISPQ TVWDYVEKIKASGTKEICVVRFTPVTEEDQISYT LLFAYFSSRKRYGVAANNMKQVKDMYLIPLGAT DKIPHPLVPFDGPGLELHRPNLLLGLIIRQKLKRQ

Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, Predicted end Predicted SEO ID Method E-Glutamic Acid, F-Phenylalanine, G-Glycine, H-Histidine, nucleotide beginning NO: I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, nucleotide location N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, corresponding location T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, corresponding to last amino X=Unknown, *=Stop codon, /=possible nucleotide deletion, acid residue of to first amino >=possible nucleotide insertion acid residue of peptide sequence peptide sequence **HSACASTSHIAETPESAPPIALPPDKKSKIEVSTEE APEEENDFFNSFTTVLHKQRNKPQQNLQEDLPTA** VEPLMEVTKQEPPKPLRFLPGVLIGWENQPTTLE LANKPLPVDDILQSLLGTTGQVYDQ\AQSVMEQ NTVKEIPFLNEOTNSKIEKTDNVEVTDGENKEIK VKVDNISESTDKSAEIETSVVGSSSISAGSLTSLSL RGKPPDVSTEAFLTNLSIQSKQEETVESKEKTLKR QLQEDQENNLQDNQTSNSSPCRSNVGKGNIDGN VSCSENLVANTARSPQFINLKRDPRQAAGRSQPV TTSESKDGDSCRNGEKHMLPGLSHNKEHLTEQIN VEEKLCSAEKNSCVQQSDNLKVAQNSPSVENIQT SOAEQAKPLQEDILMQNIETVHPFRRGSAVATSH FEVGNTCPSEFPSKSITFTSRSTSPRTSTNFSPMRP **OOPNLOHLKSSPPGFPFPGPPNFPPQSMFGFPPHL** PPPLLPPPGFG\FA\QNPMVPWPPVV\HLP\GQPQR MMGPLSQASRYIGPQNFYQVKDIRRPERRHSDP WGRQDQQLDRPFNRGKGDRQRFYSDSHHLKR ERHEKEWEQESERHRRRDRSQDKDRDRKSREEG HKDKERARLSHGDRGTDGKASRDSRNVDKKPD KPKSEDYEKDKEREKSKHREGEKDRDRYHKDR DHTDRTKSKR GSWTVELSLKPSASPSLKWVCLPGAAAVNKHRS 3539 157 1769 A GAGGLIRSLIQCTWAPAGPARRGGRGIEDFPYLF FQLTHCQQRICSVTQAGVQWCDHSSLQPQTPGL NOSSHLSLLSSRDYRMLSSFNEWFWQDRFWLPP NYTWTELEDRDGRVYPHPQDLLAALPLALVLLA MRLAFERFIGLPLSRWLGVRDQTRRQVKPNATL **EKHFLTEGHRPKEPQLSLLAAQCGLTLQQTQRW** FRRRNQDRPQLTKKFCEASWRFLFYLSSFVGGL SVLYHESWLWAPVMCWDRYPNQLTLSCPAADS EA\SLYWWYLLELGFYLSLLIRLPFDVKRKGGGP SSIKPRPHYDPPSTA\DFKEQVIHHFVAVILMTFSY SANLLRIGSLVLLLHDSSDYLLEACKMVNYMQY QQVCDALFLIFSFVFFYTRLVLFPTQILYTTYYESI SNRGPFFGYYFFNGLLMLLQLLHVFWSCLILRML YSFMKKGQMEKDIRSDVEESDSSEEAAAAQEPL OLKNGTAGGPRPAPTDGPRSRVAGRLTNRHTTA SPAGYCHSGLLPGCSRSA/CADLAKHQELPGKKL 3540 267 1397 A LSEKKLKRYFVDYRRVLVCGGNGGAGASCFHSE PRKEFGGPDGGDGGNGGHVILRVDQQVKSLSSV LSRYQGFSGEDGGSKNCFGRSGAVLYIRVPVGTL VKEGGRVVADLSCVGDEYIAALGGAGGKGNRF FLANNNRAPVTCTPGQPGQQRVLHLELKTVAHA **GMVGFPNAGKSSLLRAISNARPAVASYPFTTLKP** HVGIVHYEGHLQIAVADIPGIIRGAHQNRGLGSA FLRHIERCRFLLFVVDLSQPEPWTQVDDLKYELE MYEKGLSARPHAIVANKIDLPEAQANLSQLRDH LGQEVIVLSALTGENLEQLLLHLKVLYDAYAEA **ELGQGRQPLRW** DTQVSETLKRFAGKVTTASVKERREILSELGKCV 8008 3541 A 1 AGKDLPEGAVKGLCKLFCLTLHRYRDAASRRAL QAAIQQLAEAQPEATAKNLLHSLQSSGIGSKAGV **PSKSSGSAALLALTWTCLLVRIVFPSRAKRQGDI** WNKLVEVQCLLLLEVLGGSHKHAVDGAVKKLT

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Predicted end Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, SEQ ID Method Predicted beginning nucleotide E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, NO: nucleotide location I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, corresponding N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, location corresponding to last amino T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, acid residue of X=Unknown, *=Stop codon, /=possible nucleotide deletion, to first amino acid residue of peptide \=possible nucleotide insertion peptide sequence sequence KLWKENPGLVEQYLSAILSLEPNQNYAGMLGLL VOFCTSHKEMDVVSQHKSALLDFYMKNILMSK VKPPKYLLDSCAPLLRYLSHSEFKDLILPTIQKSL LRSPENVIETISSLLASVTLDLSQYAMDIVKGLAG HLKSNSPRLMDEAVLALRNLARQCSDSSAMESL TKHLFAILGGSEGKLTVVAQKMSVLSGIGSVSHH **VVSGPSSQVLNGIVAELFIPFLQQEVHEGTLVHA** VSVLALWCNRFTMEVPKKLTEWFKKAFSLKTST SAVRHAYLQCMLASYRGDTLLQALDLLPLLIQT VEKAASOSTOVPTITEGVAAALLLLKLSVADSQA EAKLSSFWQLIVDEKKQVFTSEKFLVMASEDAL CTVLH\LTERLFLDHPHRLTGNKVQOYHRALVA VLLSRTWHVRRQAQQTVRKLLSSLGGFKLAHGL LEELKTVLSSHKVLPLEALVTDAGEVTEAGKAY VPPRVLQEALCVISGVPGLKGDVTDTEQLAQEM LIISHHPSLVAVQSGLWPALLARMKIDPEAFITRH LDQIIPRMTTQSPLNQSSMNAMGSLSVLSPDRVL POLISTITASVONPALRLVTREEFAIMQTPAGELY DKSIIQSAQQDSIKKANMKRENKAYSFKEQIIELE LKEEIKKKKGIKEEVQLTSKQKEMLQAQLDREA QVRRRLQELDGELEAALGLLDIILAKNPSGLTQYI PVLVDSFLPLLKSPLAAPRIKNPFLSLAACVMPSR LKALGTLVSHVTLRLLKPECVLDKSWCQEELSV AVKRAVMLLHTHTITSRVGKGEPGAAPLSAPAFS LVFPFLKMVLTEMPHHSEEEEEWMAQILQILTVQ AQLRASPNTPPGRVDENGPELLPRVAMLRLLTW VIGTGSPRLQVLASDTLTTLCASSSGDDGCAFAE QEEVDVLLCALQSPCASVRETVLRGLMELHMVL PAPDTDEKNGLNLLRRLWVVKFDKEEEIRKLAE RLWSMMGLDLQPDLCSLLIDDVIYHEAAVRQAG **AEALSQAVARYQRQAAEVMGRLMEIYQEKLYR** PPPVLDALGRVISESPPDQWEARCGLALALNKLS OYLDSSQVKPLFQFFVPDALNDRHPDVRKCMLD AALATLNTHGKENVNSLLPVFEEFLKNAPNDAS YDAVRQSVVVLMGSLAKHLDKSDPKVKPIVAKL IAALSTPSQQVQESVASCLPPLVPAIKEDAGGMIQ RLMQQLLESDKYAERKGAAYGLAGLVKGLGILS LKQQEMMAALTDAIQDKKNFRRREGALFAFEM LCTMLGKLFEPYVVHVLPHLLLCFGDGNQYVRE AADDCAKAVMSNLSAHGVKLVLPSLLAALEEES WRTKAGSVELLGAMAYCAPKQLSSCLPNIVPKL TEVLTDSHVKVQKAGQQALRQIGSVIRNPEILAI APVLLDALTDPSRKTQKCLQTLLDTKFVHFIDAP SLALIMPIVQRAFQDRSTDTRKMAAQIIGNMYSL TDQKDLAPYLPSVTPGLKASLLDPVPEVRTVSAK ALGAMVKGMGESCFEDLLPWLMETLTYEQSSV DRSGAAQGLAEVMAGLGVEKLEKLMPEIVATAS KVDIAPHVRDGYIMMFNYLPITFGDKFTPYVGPII **PCILKALADENEFVRDTALRAGQRVISMYAETAI ALLLPQLEQGLFDDLWRIRFSSVQLLGDLLFHISG** VTGKMTTETASEDDNFGTAQSNKAIITALGVERR NRVLAGLYMGRSDTQLVVRQASLHVWKIVVSN **TPRTLREILPTLFGLLLGFLASTCADKRTIAARTL GDLVRKLGEKILPEIIPILEEGLRSQKSDERQGVCI** GLSEIMKSTSRDAVLYFSESLVPTARKALCDPLE

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SEQ ID	Method	Predicted	Predicted end nucleotide	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine,
NO:		beginning nucleotide	location	I=Isoleucine, K=Lysine, L=Leucine, M=Methionine,
		location	corresponding	N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine,
		corresponding	to last amino	T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine,
		to first amino	acid residue of	X=Unknown, *=Stop codon, /=possible nucleotide deletion,
	1	acid residue of peptide	peptide sequence	\≔possible nucleotide insertion
		sequence	sequence	
	 	-		EVREAAAKTFEQLHSTIGHQALEDILPFLLKQLD
]	}	}		DEEVSEFALDGLKQVMAIKSRVVLPYLVPKLTTP
]	}		PVNTRVLAFLSSVAGDALTRHLGVILPAVMLAL
i				KEKLGTPDEQLEMANCQAVILSVEDDTGHRIIIE
				DLLEATRSPEVGMRQAAAIILNIYCSRSKADYTS
				HLRSLVSGLIRLFNDSSPVVLEESWDALNAITKK
}	l		1	LDAGNQLALIEELHKEIRLIGNESKGEHVPGFCLP
	[KKGVTSILPVLREGVLTGSPEQKEEAAKALGLVI
				RLTSADALRPSVVSITGPLIRILGDRFSWNVKAAL
t I			ŀ	LETLSLLLAKVGIALKPFLPQLQTTFTKALQDSNR
ł		j]	GVRLKAADALGKLISIHIKVDPLFTELLNGIRAME
[1			DPGVRDTMLQALRFVIQGAGAKVDAVIRKNIVS
1				LLLSMLGHDEDNTRISSAGCLGELCAFLTEELS
			ļ	AVLQQCLLADVSGIDWMVRHGRSLALSVAVNV
ĺ			Ì	APGRLCAGRYSSDVQEMILSSATADRIPIAVSGV
į.			Ì	RGMGFLMRHHIETGGGQLPAKLSSLFVKCLQNP
				SSDIRLVAEKMIWWANKDPLPPLDPQAIKPILKA
Ì				LLDNTKDKNTVVRAYSDQAIVNLLKMRQGEEVF
		ļ		QSLSKILDVASLEVLNEVNRRSLKKLASQADSTE
	ļ. <u>.</u>			QVDDTILT
3542	A	62	1130	PWNPQDFPGNRGLMG\QKGEIGPP\GQQGKKGAP
				GMP\GLMGSNGSPGQPGTPGSKGSKGEPGIQGMP
		{		GASGLKGEPGATGSPGEPGYMGLPGIQGKKGDK
				GNQGEKGIQGQKGENGRQGIPGQQGIQGHHGAK
ŀ	1		}	GERGEKGEPGVRGAIGSKGESGVDGLMGPAGPK
Í	[İ	GQPGDPGPQGPPGLDGKPGREFSEQFIRQVCTDV
				IRAQLPVLLQSGRIRNCDHCLSQHGSPGIPGPPGPI
]	1 .	}	GPEGPRGLPGLPGRDGVPGLVGVPGRPGVRGLK
				GLPGRNGEKGSQGFGYPGEQGPPGPPGPEGPPGI
	}			SKEGPPGDPGLPGKDGDHGKPGIQGQPGPPGICD
				PSLCFSVIARRDPFRKGPNY
3543	A	654	194	PARSLEKMKASVVLSLLGYLVVPSGAYILGRCTV
}	}		1	AKKLHDGGLDYFERYSLENWVCLAYFESKFNPS\
				AIYENTREGYTGFGLFQMRGSDWCGDHGRNRC
		ļ		HMSCSALLNPNLEKTIKCAKTIVKGKEGMGAWP
				TWSRYCQYSDTLARWLDGCKL
3544	Α	2	1074	SCRLAAGRLAQWLLRASRSGMLRAGWLRGAAA
Ï				LALLLAARVVAAFEPITVGLAIGAASAITGYLSY
1			1	NDIYCRFAECCREERPLNASALKLDLEEKLFGQH
1	'	1	}	LATEVI\FKALTGFRNNKNPKKPLTLSLHGWAGT
1				GKNFVSQMGAENLHPKGLKSNFVHLFVSTLHFP
1			1	HEQKIKLYQDQLQKWIRGNVSACANSVFIFDEM
				DKL\HPGIIE\AIKPFLDYYEHVERVSYR\KAIFIFLS
	ļ	1	i	NAGGDLITKTALDFWRAGRKREDIQLKDLEPVL
l	1		1	SVGVFNNKHSGLWHSGLIDKNLIDYFIPFLPLEYR
ļ	1			HVKMCVRAEMRARGSAIDEDIVTRVAEEMTFFP\
				RDEKIYSDKGCKTVQSRLDFH
3545	A	3	273	SAQGRSWGRFYRQIKRHPGIIPMIGLICLGMGSA
			}	ALYLLRLALRSPDVW*SWDRKNNPEPWNRLSPN
			1	DQYKFLAVSTDYKKLKKDRPDF
3546	A	23	591	ALSTETRTPDMRRLLLVTSLVVVLLWEAGAVPA
7740	1	"	371	PKVPIKMQVKHWPSEQDPEKAWGARVVEPPEK
Ī		1		DDQLVVLFPVQKPKLLTTEEKPRGQGRGPILPGT
1	1	1		KAWMETEDTLGRVLSPEPDHDSLYHPPPEEDQG
}		1		EERPRLWVMPNHQVLLGPEEDQDHIYHPQ*GSR
L	<u></u>	<u></u>	1	EEVLYT A MATHUM A PROLECTION I ULA . 02K

			I maratar v	L Amino cold anguages (A-Alonino Ca-Cantaino N. A
SEQ ID NO:	Method	Predicted beginning nucleotide location corresponding to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion
		sequence		GHHCPRPVPRPRLLGLGPSLPCPS
3547	A	23	591	ALSTETRTPDMRRLLLVTSLVVVLLWEAGAVPA PKVPIKMQVKHWPSEQDPEKAWGARVVEPPEK DDQLVVLFPVQKPKLLTTEEKPRGQGRGPILPGT KAWMETEDTLGRVLSPEPDHDSLYHPPPEEDQG EERPRLWVMPNHQVLLGPEEDQDHIYHPQ*GSR GHHCPRPVPRPRLLGLGPSLPCPS
3548	A		1641	TWLPSVPAEEVQQPEMAAVLNAERLEVSVDGLT LSPDPEERPGAEGAPLAAATAATALATWIRSRPG RLRGTARSPGRRAAGGAAEEARRLEQRWGFGLE ELYGLALRFFKEKDGKAFHPTYEEKLKLVALHK QVLMGPYNPDTCPEVGFFDVLGNDRRREWAAL GNMSKEDAMVEFVKLLNRCCHLFSTYVASHKIE KEEQEKKRKEEEERRREEEERERLQKEEEKRR EEEERLRREEEERRRIEEERLRLEQQKQQIMAAL NSQTAVQFQQYAAQQYPGNYEQQQILIRQLQEQ HYQQYMQQLYQVQLAQQQAALQKQQEVVVAG SSLPTSSKVECNCTQVI*CQFNRQAKTHTDSSEKE LEPEAAEEALENGPKESLPVIAAPSMWTRPQIKD FKEKIQQDADSVITVGRGEVVTVRVPTHEEGSYL FWEFATDNYDIGFGVYFEWTDSPNTAVSVHVSE SSDDDEEEEENIGCEEKAKKNANKPLLDEIVPVY RRDCHEEVYAGSHQYPGRGVYLLKFDNSYSLW RSKSVYYRVYYTR
3549	A	1837	3593	PAVLVLEPASQSRKQQNTASATAQHWSAQIHKE SFLAPVFTKDEQKHRRPYEFEVERDAKARGLEQF SATHGHTPIILNGWHGESAMDLSCSSEGSPGATS PFPVSASTPKIGAISSLQGALGMDLSGILQAGLIHP VTGQIVNGSLRRDDAATRRRGRRKHVEGGMD LIFLKEQTLQAGILEVHEDPGQATLSTTHPEGPGP ATSAPEPATAASSQAEKSIPSKSLLDWLRQQADY SLEVPGFGANFSDKPKQRRPRCKEPGKLDVSSLS GEERVPAIPKEPGLRGFLPENKFNHTLAEPILRDT GPRRRGRRPRSELLKAPSIVADSPSGMGPLFMNG LIAGMDLVGLQNMRNMPGIPLTGLVGFPAGFAT MPTGEEVKSTLSMLPMMLPGMAAVPQMFGVGG LLSPPMATTCTSTAPASLSSTTKSGTAVTEKTAE DKPSSHDVKTDTLAEDKPGPGPFSDQSEPAITTSS PVAFNPFLIPGVSPGLIYPSMFLSPGMGMALPAM QQARHSEIVGLESQKRKKKKTKGDNPNSHPEPA PSCEREPSGDENCAEPSAPLPAEREHGAQAGEGA LKDSNNDTN
3550	A	287	39	QLNLNKIATSQKHRDFVAESVGEKPVGSLAGIGE VMDKKLEEGCFDKAYVVLGQFLVLKKDEDLF*E WLRDTGGARTRGSRE
3551	A	21	3925	GDLLEVGLPPGLEFPRGICLRGLRRTMSLDFGSV ALPVQNEDEEYDEEDYEREKELQQLLTDLPHDM LDDDLSSPELQYSDCSEDGTDGQPHHPEQLEMS WNEQMLPKSQSVNGPSCQGLEPYNKVTYKPYQS SAQNNGSPAQEITGSDTFEGLQQQFLGANENSAE NMQIIQLQVLNKAKERQLENLIEKLNESERQIRY LNHQLVIIKDEKDGLTLSLRESQKLFQNGKEREIQ LEAQIKALETQIQALKVNEEQMIKKSRTTEMALE SLKQQLVDLHHSESLQRAREQHESIVMGLTKKY EEQVLSLQKNLDATVTALKEQEDICSRLKDHVK

SEQ ID NO:	Method	Predicted beginning nucleotide location corresponding to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion
				QLERNQEAIKLEKTEIINKLTRSLEESQKQCAHLL QSGSVQEVAQLQFQLQQAQKAHAMSANMNKA LQEELTELKDEISLYESAAKLGIHPSDSEGELNIEL TESYVDLGIKKVNWKKSKVTSIVQEEDPNEELSK DEFILKLKAEVQRLLGSNSMKRHLVSQLQNDLK DCHKKIEDLHQVKKDEKSIEVETKTDTSEKPKNQ LWPESSTSDVVRDDILLLKNEIQVLQQQNQELKE TEGKLRNTNQDLCNQMRQMVQDFDHDKQEAV DRCERTYQQHHEAMKTQIRESLLAKHALEKQQL FEAYERTHLQLRSELDKLNKEVTAVQECYLEVC REKDNLELTLRKTTEKEQQTQEKIKEKLIQQLEK EWQSKLDQTIKAMKKKTLDCGSQTDQVTTSDVI SKKEMAIMIEEQKCTIQQNLEQEKDIAIKGAMKK LEIELELKHCENITKQVEIAVQNAHQRWLGELPE LAEYQALVKAEQKKWEEQHEVSVNKRISFAVSE AKEKWKSELENMRKNILPGKELEEKIHSLQKELE LKNEEVPVVIRAELAKARSEWNKEKQEEIHRIQE QNEQDYRQFLDDHRNKINEVLAAAKEDFMKQK TELLLQKETELQTCLDQSRREWTMQEAKRIQLEI YQYEEDILTVLGVLLSDTQKEHISDSEDKQLLEI MSTCSSKWMSVQYFEKLKGCIQKAFQDTLPLLV ENADPEWKKRNMAELSKDSASQGTGQGDPGPA AGHHAQPLALQATEAEADKKKVLEIKDLCCGHC FQELEKAKQECQDLKGKLEKCCRHLQHLERKHK AVVEKIGEENNKVVEELIEENNDMKNKLEELQT LCKTPPRSLSAGAIENACLPCSGGALEELRGQYIK AVKKIKCDMLRYIQESKERAAEMVKAEVL*ERQ
3552	A	771	375	ETARKMRKYYLICLQQILQDDGKEGAEKKIMNA ASKLATMAKLLETPISSKSQSKTTQSGMSK ARTRQTSGQAREPEKESPAPGGGGLAEIRSRQQL SQTSRIPPLAKDQAVEAMFPPARGKELLSFEDVA
				MYFTREEWGHLNWGQKDLYRDVMLENYRNMV LLVYFQFDAAIPLC*TSLAHSSWLQLYFRLYF
3553	A	76	72	PGVRGVEAPGGVAPGRNAMRRGERRDAGGPRP ESPVPAGRASLEEPPDGPSAGQATGPGEGRRSTE SEVYDDGTNTFFWRAHTLTVLFILTCTLGYVTLL EETPQDTAYNTKRGIVASILVFLCFGVTQAKDGP FSRPHPAYWRFWLCVSVVYELFLIFILFQTVQDG RQFLKYVDPKLGVPLPERDYGGNCLIYDPDNET DPFHNIWDKLDGFVPAHFLGWYLKTLMIRDWW MCMIISVMFEFLEYSLEHQLPNFSECWWDHWIM DVLVCNGLGIYCGMKTLEWLSLKTYKWQGLWN IPTYKGKMKRIAFQFTPYSWVRFEWKPASSLRR WLAVCGIILVFLLAELNTFYLKFVLWMPPEHYLV LLRLVFFVNVGGVAMREIYDFMDDPKPHKKLGP QAWLVAAITATELLIVVKYDPHTLTLSLPFYISQC WTLGSVLALTWTVWRFFLRDITLRYKETRWQK WQNKDDQGSTVGNGDQHPLGLDEDLLGPGVAE GEGAPTPN*PRGPAPRPLPSAPRAVCGASSRR
3554	A	2	2106	FDEFSALPSPSLQTSWSFGPMSRRALRRLRGEQR GQEPLGPGALHFDLRDDDDAEEEGPKRELGVRR PGGAGKEGVRVNNRFELINIDDLEDDPVVNGERS GCALTDAVAPGNKGRGQRGNTESKTDGDDTET VPSEQSHASGKLRKKKKKQKNKKSSTGEASENG LEDIDRILERIEDSTGLNRPGPAPLSSRKHVLYVE

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SEQ ID NO:	Method	Predicted beginning nucleotide location corresponding to first amino acid residue of peptide	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion
		sequence		HRHLNPDTELKRYFGARAILGEQRPRQRQRVYP KCTWLTTPKSTWPRYSKPGLSMRLLESKKGLSFF AFEHSEEYQQAQHKFLVAVESMEPNNIVVLLQT SPYHVDSLLQLSDACRFQEDQEMARDLVERALY SMECAFHPLFSLTSGACRLDYRRPENRSFYLALY KQMSFLEKRGCPRTALEYCKLILSLEPDEDPLCM LLLIDHLALRARNYEYLIRLFQEWEVGASLAHRN LSQLPNFAFSVPLAYFLLSQQTDLPECEQSSARQ KASLLIQQALTMFPGVLLPLLESCSVRPDASVSSH RFFGPNAEISQPPALSQLVNLYLGRSHFLWKEPA TMSWLEENVHEVLQAVDAGDPAVEACENRRKV LYQRAPRNIHRHVILSEIKEAVAALPPDVTTQSV MGFDPLPPSDTIYSYVRPERLSPISHGNTIALFFRS LLPNYTMEGERPEEGVAGGLNRNQGLNRLMLA VRDMMANFHLNDLEAPHEDDA*GEGEWD
3555	A	2	2106	FDEFSALPSPSLQTSWSFGPMSRRALRRLRGEQR GQEPLGPGALHFDLRDDDDAEEEGPKRELGVRR PGGAGKEGVRVNNRFELINIDDLEDDPVVNGERS GCALTDAVAPGNKGRGQRGNTESKTDGDDTET VPSEQSHASGKLRKKKKKKQKNKKSSTGEASENG LEDIDRILERIEDSTGLNRPGPAPLSSRKHVLYVE HRHLNPDTELKRYFGARAILGEQRPRQRQRVYP KCTWLTTPKSTWPRYSKPGLSMRLLESKKGLSFF AFEHSEEYQQAQHKFLVAVESMEPNNIVVLLQT SPYHVDSLLQLSDACRFQEDQEMARDLVERALY SMECAFHPLFSLTSGACRLDYRRPENRSFYLALY KQMSFLEKRGCPRTALEYCKLILSLEPDEDPLCM LLLIDHLALRARNYEYLIRLFQEWEVGASLAHRN LSQLPNFAFSVPLAYFLLSQQTDLPECEQSSARQ KASLLIQQALTMFPGVLLPLLESCSVRPDASVSSH RFFGPNAEISQPPALSQLVNLYLGRSHFLWKEPA TMSWLEENVHEVLQAVDAGDPAVEACENRRKV LYQRAPRNIHRHVILSEIKEAVAALPPDVTTQSV MGFDPLPPSDTIYSYVRPERLSPISHGNTIALFFRS LLPNYTMEGERPEEGVAGGLNRNQGLNRLMLA VRDMMANFHLNDLEAPHEDDA*GEGEWD
3556	A	3388	1650	KTRGTMFYYPNVLQRHTGCFATIWLAATRGSRL VKREYLRVNVVKTCEEILNYVLVRVQPPQPGLP RPRFSLYLSAQLQIGVIRVYSQQCQYLVEDIQHIL ERLHRAQLQIRIDMETELPSLLLPNHLAMMETLE DAPDPFFGMMSVDPRLPSPFDIPQIRHLLEAAIPE RVEEIPPEVPTEPREPERIPVTVLPPEAITILEAEPIR MLEIEGERELPEVSRRELDLLIAEEEEAILLEIPRL PPPAPAE*GQELLDQVGCQCWEGSPHFSCPFPLR VEGMGEALGPEELRLTGWEPGALLMEVTPPEEL RLPAPPSPERRPPVPPPPRRRRRRRLLFWDKETQI SPEKFQEQLQTRAHCWECPMVQPPERTIRGPAEL FRTPTLSGWLPPELLGLWTHCAQPPPKALRRELP EEAAAEEERRKIEVPSEIEVPREALEPSVPLMVSL EISLEAAEEEKSRISLIPPEERWAWPEVEAPEAPA LPVVPELPEVPMEMPLVLPPELELLSLEAVHRAV ALELQANREPDFSSLVSPLSPRRMAARVFYLLLV LSAQQILHVKQEKPYGRLLIQPGPRFH
3557	A	3388	1650	KTRGTMFYYPNVLQRHTGCFATIWLAATRGSRL VKREYLRVNVVKTCEEILNYVLVRVQPPQPGLP

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SEQ ID NO:	Method	Predicted beginning	Predicted end nucleotide	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine,
1.0.		nucleotide	location	I=Isoleucine, K=Lysine, L=Leucine, M=Methionine,
	1	location	corresponding	N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine,
		corresponding	to last amino	T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine,
	ĺ	to first amino acid residue of	acid residue of peptide	X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion
		peptide	sequence	- Possible nucleonae inser aou
		sequence		
				RPRFSLYLSAQLQIGVIRVYSQQCQYLVEDIQHIL
	}]]	ERLHRAQLQIRIDMETELPSLLLPNHLAMMETLE
				DAPDPFFGMMSVDPRLPSPFDIPQIRHLLEAAIPE
ļ				RVEEIPPEVPTEPREPERIPVTVLPPEAITILEAEPIR
ļ			}	MLEIEGERELPEVSRRELDLLIAEEEEAILLEIPRL
Ì				PPPAPAE*GQELLDQVGCQCWEGSPHFSCPFPLR
]		J	VEGMGEALGPEELRLTGWEPGALLMEVTPPEEL
1				RLPAPPSPERRPPVPPPPRRRRRRRLLFWDKETQI
ļ				SPEKFQEQLQTRAHCWECPMVQPPERTIRGPAEL
1				FRTPTLSGWLPPELLGLWTHCAQPPPKALRRELP
				EEAAAEEERRKIEVPSEIEVPREALEPSVPLMVSL
			ł	EISLEAAEEEKSRISLIPPEERWAWPEVEAPEAPA
				LPVVPELPEVPMEMPLVLPPELELLSLEAVHRAV
}	ļ]	ALELQANREPDFSSLVSPLSPRRMAARVFYLLLV
				LSAQQILHVKQEKPYGRLLIQPGPRFH
3558	A	489	2360	IRPRPRGRRRALDSPNAAAPPVYVCRSPGEPTSL
				VNMASEDIAKLAETLAKTQVAGGQLSFKGKSLK
,				LNTAEDAKDVIKEIEDFDSLEALRLEGNTVGVEA
l				ARVIAKAL*KKSELKRCHWSDMFTGRLRTEIPPA
	ļ		{ .	LISLGEGLITAGAQLVELDLSDNAFGPDGVQGFE
				ALLKSSACFTLQELKLNNCGMGIGGGKILAAALT
				ECHRKSSAQGKPLALKVFVAGRNRLENDGATAL
			[AEAFRVIGTLEEVHMPQNGINHPGITALAQAFAV
}				NPLLRVINLNDNTFTEKGAVAMAETLKTLRQVE
ł			•	VINFGDCLVRSKGAVAIADAIRGGLPKLKELNLS
1			· ·	FCEIKRDAALAVAEAMADKAELEKLDLNGNTLG
ł				EEGCEQLQEVLEGFNMAKVLASLSDDEDEEEEE
				EGEEEEEAEEEEEDEEEEEEEEEEEPQQRG
İ			ļ	QGEKSATPSRKILDPNTGEPAPVLSSPPPADVSTF
Ī			<u> </u>	LAFPSPEKLLRLGPKSSVLIAQQTDTSDPEKVVSA
ļ			}	FLKVSSVFKDEATVRMAVQDAVDALMQKAFNS
	1			SSFNSNTFLTRLLVHMGLLKSEDKVKAIANLYGP
ł	-	}		LMALNHMVQQDYFPKALAPLLLAFVTKPNSALE
				SCSFARHSLLQTLYKV
3559	A	489	2360	IRPRPRGRRRALDSPNAAAPPVYVCRSPGEPTSL
	ł			VNMASEDIAKLAETLAKTQVAGGQLSFKGKSLK
				LNTAEDAKDVIKEIEDFDSLEALRLEGNTVGVEA
	ľ			ARVIAKAL*KKSELKRCHWSDMFTGRLRTEIPPA
		ļ	İ	LISLGEGLITAGAQLVELDLSDNAFGPDGVQGFE
				ALLKSSACFTLQELKLNNCGMGIGGGKILAAALT
				ECHRKSSAQGKPLALKVFVAGRNRLENDGATAL
		İ	1	AEAFRVIGTLEEVHMPQNGINHPGITALAQAFAV
1	ĺ		1	NPLLRVINLNDNTFTEKGAVAMAETLKTLRQVE
		ļ		VINFGDCLVRSKGAVAIADAIRGGLPKLKELNLS
		1		FCEIKRDAALAVAEAMADKAELEKLDLNGNTLG
	1	1	ļ	EEGCEQLQEVLEGFNMAKVLASLSDDEDEEEEE
		1		EGEEEEEAEEEEEDEEEEEEEEEEPQQRG
1	1	1		QGEKSATPSRKILDPNTGEPAPVLSSPPPADVSTF
		1		LAFPSPEKLLRLGPKSSVLIAQQTDTSDPEKVVSA
		1		FLKVSSVFKDEATVRMAVQDAVDALMQKAFNS
j		1	}	SSFNSNTFLTRLLVHMGLLKSEDKVKAIANLYGP
1		1	Į	LMALNHMVQQDYFPKALAPLLLAFVTKPNSALE
L				SCSFARHSLLQTLYKV
3560	A	2	1198	FVRELPRPRPGAATAAIMVSVINTVDTSHEDMIH
				DAQMDYYGTRLATCSSDRSVKIFDVRNGGQILIA
				

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SEQ ID NO:	Method	Predicted beginning nucleotide location corresponding to first amino acid residue of peptide	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, _=possible nucleotide insertion
		sequence		DLRGHEGPVWQVAWAHPMYGNILASCSYDRKV IIWREENGTWEKSHEHAGHDSSVNSVCWAPHDY GLILACGSSDGAISLLTYTGEGQWEVKKINNAHT
				IGCNAVSWAPAVVPGSLIDHPSGQKPNYIKRFAS GGCDNLIKLWKEEEDGQWKEEQKLEAHSDWVR DVAWAPSIGLPTSTIASCSQDGRVFIWTCDDASS NTWSPKLLHKFNDVVWHVSWSITANILAVSGGD
				NKVTLWKESVDGQWVCISDVNKGQGSVSASVT EGQQNEQ*QDRWGLAPHPPAPGLPLPGPTNQTT GKSPQLQQDYFPRRSYRCSHRLIICLNVIGDAL
3561	A	540	86	WRVKEMTSTLPKALGRKTASRSHTTLQGGSCCP VLWTAKLRCRKLRFPLPPPPPSSSAWPWQGWGI RGEQEAEGPLGETGPPVGPELSGLRQWRKLIKGR YGEWRGSGQKTGQPS*TTMQGGETEENRTETTT GNKQRESEAPWVRHTYIT
3562	A	1920	242	PMMAMPFFERFKSSIQRPSPVLVLSQNTKRESGR KVQSGNINAAKTIADIIRTCLGPKSMMKMLLDP MGGIVMTNDGNAILREIQVQHPAAKSMIEISRTQ
				DEEVGDGTTSVIILAGEMLSVAEHFLEQQMHPTV VISAYRKALDDMISTLKKISIPVDISDSDMMLNIIN SSITTKAISRWSSLACNIALDAVKMVQFEENGRK EIDIKKYARVEKIPGGIIEDSCVLRGVMINKDVTH PRMRRYIKNPRIVLLDSSLEYKKGESQTDIEITRE
÷ .	 - 			EDFTRILQMEEEYIQQLCEDIIQLKPDVVITEKGIS DLAQHYLMRANITAIRRVRKTDNNRIARACGARI VSRPEELREDDVGTGAGLLEIKKIGDEYFTFITDC KDPKACTILLRGASKEILSEVERNFQDAMQVCRN
				VLLDPQLVPGGGASEMAVAHALTEKSKAMTGV EQWPYRAVAQALEVIPRTLIQNCGASTIRLLTSLR AKHTQENCETWGVNGETGTLVDMKELGIWEPL AVKLQTYKTAVETAVLLLRIDDIVSGHKKKGDD QSRQGGAPDAGQE
3563	A	1571	560	GPSLLGTRGTPNPARTLQIFFLIIGRRLTGRMAAV DDLQFEEFGNAATSLTANPDATTVNIEDPGETPK
				HQPGSPRGSGREEDDELLGNDDSDKTELLAGQK KSSPFWTFEYYQTFFDVDTYQVFDRIKGSLLPIPG KNFVRLYIRSNPDLYGPFWICATLVFAIAISGNLS NFLIHLGEKTYHYVPEFRKVSIAATIIYAYAWLVP
				LALWGFLMWRNSKVMNIVSYSFLEIVCVYGYSL FIYIPTAILWIIPHKAVRWILVMIALGISGSLLAMT FWPAVREDNRRVALATIVTIVLLHMLLSVGCLA YFFDAPEMDHLPTTTATPNQTVAAAKSS
3564	A	1	328	NSRVDDFVAHLQRPLLGPASCLGILRPAMTAHSF ALPGIIFTTFWGLVGIAGPWFVPKGPNRGVIITML VATAVCCYLFWLIAILAQLNPLFGPQLKNETIWY VRFLWE
3565	A	2	1081	FVTDFPARSMAATSLMSALAARLLQPAHSCSLRL RPFHLAAVRNEAVVISGRKLAQQIKQEVRQEVEE WVASGNKRPHLSVILVGENPASHSYVLNKTRAA AVVGINSETIMKPASISEEELLNLINKLNNDDNVD GLLVQLPLPEHIDERRICNAVSPDKDVDGFHVIN VGRMCLDQYSMLPATPWGVWEIIKRTGIPTLGK NVVVAGRSKNVGMPIAMLLHTDGAHERPGGDA TVTISHRYTPKEQLKKHTILADIVISAAGIPNLITA DMIKEGAAVIDVGINRVHDPVTAKPKLVGDVDF

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SEQ ID NO:	Method	Predicted beginning nucleotide location corresponding to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \+=possible nucleotide insertion
				EGVRQKAGYITPVPGGVGPMTVAMLMKNTIIAA KKVLRLEEREVLKSKELGVATN
3566	A	3	1130	SCRRGRQQRRNVSLSSQFAHTMAAPAQQTTQP GGGKRKGKAQYVLAKRARRCDAGGPRQLEPGL QGILITCNMNERKCVEEAYSLLNEYGDDMYGPE KFTDKDQQPSGSEGEDDDAEAALKKEVGDIKAS TEMRLRRFQSVESGANNVVFIRTLGIEPEKLVHHI LQDMYKTKKKKTRVILRMLPISGTCKAFLEDMK KYAETFLEPWFKAPNKGTFQIVYKSRNNSHVNR EEVIRELAGIVCTLNSENKVDLTNPQYTVVVEIIK AVCCLSVVKDYMLFRKYNLQEVVKSPKDPSQLN SKQGNGKEAKLESADKSDQNNTAEGKNNQQVP ENTEELGQTKPTSNPQVVNEGGAKPELASQATE GSKSNENDFS
3567	A	248	3498	GKKDSSPWTCPFHPPLQLFFVIRNTRQLGDFHLA KIKVRNYWTADGDLDIGAKNVKLYVNRNLIFNG KLDKGDREAPADHSILVDQKNEKSEQLEEAMNA HSEESKGTHEMAGASGDKELGLGCSPPAETLAD AKLSSQGNVSGKRKNSTNCRKDSLSQLEEYLRLS AVPTSMGDMPSAPATSPPVKCPPVHEEPSLIQQL ENLMGRKICEPPGKTPSWLQPSPTGKDRKQGGR KPKPLWLSPEKPLAWKGRLPSDDVIGEGPGETEA RDKGLRHEPGWGTSRSVNTKERPQRATTKVHSD DSDIFNQPPNRERPASGRRGSRKDAGSSSHGDDQ
			-	PASREDTWSSRTPSRSRWRSEQEHTLHESWSSLS AFDRSHRGRISNTELPGDILDELLQQKSSRHSDLP PSKKGEQPGLSRGQDGYSGETDAGGDFKIPVLPY GQRLVIDIKSTWGDRHYVGLNGIEIFSSKGEPVQI SNIKADPPDINILPAYGKDPRVVTNLIDGVNRTQ DDMHVWLAPFTRGRSHSITIDFTHPCHVALIRIW NYNKSRIHSFRGVKDITMLLDTQCIFEGEIAKASG TLAGAPEHFGDTILFTTDDDILEAIFYSDEMFDLD VGSLDSLQDEEAMRRPSTADGEGDERPFTQAGL GADERIPELELPSSSPVPQVTTPEPGIYHGICLQLN FTASWGDLHYLGLTGLEVVGKEGQALPIHLHQIS ASPRDLNELPEYSDDSRTLDKLIDGTNITMEDEH MWLIPFSPGLDHVVTIRLDRAESIAGLRFWNYNK SPEDTYRGAKIVHVSLDGLCVSPPEGFLIRKGPG NCHFDFAQEILFVDYLRAQLLPQPARRLDMRSLE CASMDYEAPLMPCGFIFQFQLLTSWGDPYYIGLT GLELYDERGEKIPLSENNIAAFPDSVNSLEGVGG DVRTPDKLIDQVNDTSDGRHMWLAPILPGLVNR VYVIFDLPTTVSMIKLWNYAKTPHRGVKEFGLL VDDLLVYNGILAMVSHLVGGILPTCEPTVPYHTI LFTEDRDIRHQEKHTTISNQAEDQDVQMMNENQ IITNAKRKQSVVDPALRPKTCISEKETRRRC
3568	A .	50	1724	AQGGTLSAASRFCRGGLLGPWLHPASEMAATLD LKSKEEKDAELDKRIEALRRKNEALIRRYQEIEE DRKKAELEGVAVTAPRKGRSVEKENVAVESEKN LGPSRRSPGTPRPPGASKGGRTPPQQGGRAGMG RASRSWEGSPGEQPRGGGAGGRGRRGRGRGSPH LSGAGDTSISDRKSKEWEERRRQNIEKMNEEME KIAEYERNQREGVLEPNPVRNFLDDPRRRSGPLE ESERDRREESRRHGRNWGGPDFERVRCGLEHER QGRRAGLGSAGDMTLSMTGRERSEYLRWKQER

	N1	D31-4-3	Dundint	Amine said sequence (A-Alemine C-Custoine D-Assentis A-13
SEQ ID NO:	Method	Predicted beginning nucleotide location corresponding to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion
				EKIDQERLQRHRKPTGQWRREWDAEKTDGMFK DGPVPAHEPSHRYDDQAWARPPKPPTFGEFLSQ HKAEASSRRRKSSRPQAKAAPRAYSDHDDRWE TKEGAASPAPETPQPTSPETSPKETPMQPPEIPAP AHRPPEDEGEENEGEEDEEWEDISEDEEEEEIEVE EGDEEEPAQDHQAPEAAPTGIPCSEQAHGVPFSP EEPLLEPQAPGTPSSPFSPPSGHQPVSDWGEEVEL NSPRTTHLAGALSPGEAWPFESV
3569	A	1	912	MGRVGRAGVQLGRRRTTWAAERTGQAAAGGP GRALRGQRPDLRSGGAADSPAAGRGELYCGVLP RSPWFLSERRRQMADFDTYDDRAYSSFGGGRGS RGSAGGHGSRSQKELPTEPPYTAYVGNLPFNTV QGDIDAIFKDLSIRSVRLVRDKDTDKFKGFCYVE FDEVDSLKEALTYDGALLGDRSLRVDIAEGRKQ DKGGFGFRKGGPDDRGFRDDFLGGRGGSRPGDR RTGPPMGSRFRDGPPLRGSNMDFREPTEEERAQR PRLQLKPRTVATPLNQVANPNSAIFGGARPREEV VQKEQE
3570	A	1	912	MGRVGRAGVQLGRRRTTWAAERTGQAAAGGP GRALRGQRPDLRSGGAADSPAAGRGELYCGVLP RSPWFLSERRRQMADFDTYDDRAYSSFGGGRGS RGSAGGHGSRSQKELPTEPPYTAYVGNLPFNTV QGDIDAIFKDLSIRSVRLVRDKDTDKFKGFCYVE FDEVDSLKEALTYDGALLGDRSLRVDIAEGRKQ DKGGFGFRKGGPDDRGFRDDFLGGRGGSRPGDR RTGPPMGSRFRDGPPLRGSNMDFREPTEEERAQR PRLQLKPRTVATPLNQVANPNSAIFGGARPREEV VQKEQE
3571	A	28	131	RHFFGNLCAMRAKWRKKRMRRLKRKRRKMRQ RSK
3572	A	3	1202	QSEPHRKVRVDPPVRDRPPPHPPPLLVQRALPGQ GQAEGSDGADGAKRRAMAHQTGIHATEELKEFF AKARAGSVRLIKVVIEDEQLVLGASQEPVGRWD QDYDRAVLPLLDAQQPCYLLYRLDSQNAQGFE WLFLAWSPDNSPVRLKMLYAATRATVKKEFGG GHIKDELFGTVKDDLSFAGYQKHLSSCAAPAPLT SAERELQQIRINEVKTEISVESKHQTLQGLAFPLQ PEAQRALQQLKQKMVNYIQMKLDLERETIELVH TEPTDVAQLPSRVPRDAARYHFFLYKHTHEGDP LESVVFIYSMPGYKCSIKERMLYSSCKSRLLDSV EQDFHLEIAKKIEIGDGAELTAEFLYDEVHPKQH AFKQAFAKPKGPGGKRGHKRLIRGPGENGDDS
3573	A		1869	PHCEPNPGAGAMVLLHVLFEHAVGYALLALKEV EEISLLQPQVEESVLNLGKFHSIVRLVAFCPFASS QVALENANAVSEGVVHEDLRLLLETHLPSKKKK VLLGVGDPKIGAAIQEELGYNCQTGGVIAEILRG VRLHFHNLVKGLTDLSACKAQLGLGHSYSRAKV KFNVNRVDNMIIQSISLLDQLDKDINTFSMRVRE WYGYHFPELVKIINDNATYCRLAQFIGNRRELNE DKLEKLEELTMDGAKAKAILDASRSSMGMDISAI DLINIESFSSRVVSLSEYRQSLHTYLRSKMSQVAP SLSALIGEAVGARLIAHAGSLTNLAKYPASTVQIL GAEKALFRALKTRGNTPKYGLIFHSTFIGRAAAK NKGRISRYLANKCSIASRIDCFSEVPTSVFGEKLR EQVEERLSFYETGEIPRKNLDVMKEAMVQAEAE

Predicted end Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, SEQ ID Method Predicted NO: beginning nucleotide E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, nucleotide location corresponding N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, location corresponding to last amino T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, acid residue of X=Unknown, *=Stop codon, /=possible nucleotide deletion, to first amino acid residue of peptide \=possible nucleotide insertion peptide sequence sequence **EAAAEITRKLEKQEKKRLKKEKKRLAALALASS** ENSSSTPEÈCEETSEKPKKKKKQKPQEVPQENGM **EDPSISFSKPKKKKSFSKEELMSSDLEETAGSTSIP** KRKKSTPKEETVNDPEEAGHRSRSKKKRKFSKEE PVSSGPEEAVGKSSSKKKKKFHKASQED 2032 3574 À 284 **CGNERTARLWVQPVVSTMPQASEHRLGRTREPP** VNIQPRVGSKLPFAPRARSKERRNPASGPNPMLR PLPPRPGLPDERLKKLELGRGRTSGPRPRGPLRA DHGVPLPGSPPPTVALPLPSRTNLARSKSVSSGDL RPMGIALGGHRGTGELGAALSRLALRPEPPTLRR STSLRRLGGFPGPPTLFSIRTEPPASHGSFHMISAR SSEPFYSDDKMAHHTLLLGSGHVGLRNLGNTCF LNAVLQCLSSTRPLRDFCLRRDFRQEVPGGGRA QELTEAFADVIGALWHPDSCEAVNPTRFRAVFQ KYVPSFSGYSQQDAQEFLKLLMERLHLEINRRGR RAPPILANGPVPSPPRRGGALLEEPELSDDDRANL MWKRYLEREDSKIVDLFVGQLKSCLKCQACGY RSTTFEVFCDLSLPIPKKGFAGGKVSLRDCFNLFT KEEELESENAPVCDRCRQKTRSTKKLTVQRFPRI LVLHLNRFSASRGSIKKSSVGVDFPLQRLSLGDF ASDKAGSPVYQLYALCNHSGSVHYGHYTALCR CQTGWHVYNDSRVSPVSENQVASSEGYVLFYQL **MQEPPRCL** 3575 2408 RELDSLADLPERIKPPYANGLSTSHLRSSSVEDVK 1 Α LIISEGRPTIEVRRCSMPSVICEHTKQFQTISEESN QGSLLTVPGDTSPSPKPEVFSNVPERDLSNVSNIH SSFATSPTGASNSKYVSADRNLIKNTAPVNTVMD SPVHLEPSSQVGVIQNKSWEMPVDRLETLSTRDF ICPNSNIPDQESSLQSFCNSENKVLKENADFLSLR **QTELPGNSCAQDPASFMPPQQPCSFPSQSLSDAES** ISKHMSLSYVANQEPGILQQKNAVQIISSALDTD NESTKDTENTFVLGDVQKTDAFVPVYSDSTIQEA SPNFEKAYTLPVLPSEKDFNGSDASTQLNTHYAF SKLTYKSSSGHEVENSTTDTQVISHEKENKLESL VLTHLSRCDSDLCEMNAGMPKGNLNEODPKHC PESEKCLLSIEDEESQQSILSSLENHSQQSTQPEM HKYGQLVKVELEENAEDDKTENQIPQRMTRNK ANTMANQSKQILASCTLLSEKDSESSSPRGRIRLT **EDDDPQIHHPRKRKVSRVPQPVQVSPSLLQAKEK** TQQSLAAIVDSLKLDEIQPYSSERANPYFEYLHIR KKIEEKRKLLCSVIPQAPQYYDEYVTFNGSYLLD GNPLSKICIPTITPPPSLSDPLKELFRQQEVVRMKL RLQHSIEREKLIVSNEQEVLRVHYRAARTLANQT LPFSACTVLLDAEVYNVPLDSQSDDSKTSVRDRF NARQFMSWLQDVDDKFDKLKTCLLMRQQHEA AALNAVQRLEWQLKLQELDPATYKSISIYEIQEF YVPLVDVNDDFELTPI 3576 1421 À 5 LRLAWHDGARWPLGTPRAAATRREAAALPPVT LALLCLDGVFLSSAENDFVHRIQEELDRFLLQKQ LSKVLLFPPLSSRLRYLIHRTAENFDLLSSFSVGE **GWKRRTVICHQDIRVPSSDGLSGPCRAPASCPSR** YHGPRPISNQGAAAVPRGARAGRWYRGRKPDO PLYVPRVLRRQEEWGLTSTSVLKREAPAGRDPEE **PGDVGAGDPNSDQGLPVLMTQGTEDLKGPGQR** CENEPLLDPVGPEPLGPESQSGKGDMVEMATRF

		T-50	T 100 100 100 100 100 100 100 100 100 10	(A) (A) (A) (A) (A) (A) (A) (A) (A) (A)
SEQ ID NO:	Method	Predicted beginning nucleotide location corresponding to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion
				GSTLQLDLEKGKESLLEKRLVAEEEEDEEEVEED GPSSCSEDDYSELLQEITDNLTKKEIQIEKIHLDTS SFMEELPGEKDLAHVVEIYDFEPALKTEDLLATF SEFQEKGFRIQWVDDTHALGIFPCRASAAEALTR EFSVLKIRPLTQGTKQSKLKALQRPKLLRLVKER PQTNATVARRLVARALGLQHKKKERPAVRGPLP P
3577	A	102	1998	DTRTPGSLEMGPLQFRDVAIEFSLEEWHCLDTAQ RNLYRNVMLENYSNLVFLGIVVSKPDLIAHLEQG KKPLTMKRHEMVANPSGPVICSHFAQDLWPEQN IKDSFQKVILRRYEKRGHGNLQLIKRCESVDECK VHTGGYNGLNQCSTTTQSKVFQCDKYGKVFHK FSNSNRHNIRHTEKKPFKCIECGKAFNQFSTLITH KKIHTGEKPYICEECGKAFKYSSALNTHKRIHTG EKPYKCDKCDKAFIASSTLSKHEIIHTGKKPYKCE ECGKAFNQSSTLTKHKKIHTGEKPYKCEECGKAF NQSSTLTKHKKIHTGEKPYVCEECGKAFKYSRIL TTHKRIHTGEKPYKCNKCGKAFIASSTLSRHEFIH MGKKHYKCEECGKAFIWSSVLTRHKRVHTGEKP YKCEECGKAFKYSSTLSSHKRSHTGEKPYKCEEC GKAFVASSTLSKHEIIHTGKKPYKCEECGKAFNQ SSSLTKHKKIHTGEKPYKCEECGKAFNQSSSLTK HKKIHTGEKPYKCEECGKAFNQSSTLIKHKKIHT REKPYKCEECGKAFHLSTHLTTHKILHTGEKPYR CRECGKAFNHSATLSSHKKIHSGEKPYECDKCG KAFISPSSLSRHEIIHTGEKP
3578	A	1725	445	RPRRGTHHFSCVLGSFRVSAMFPRVSTFLPLRP LSRHPLSSGSPETSAAAIMLLTVRHGTVRYRSSA LLARTKNNIQRYFGTNSVICSKKDKQSVRTEETS KETSESQDSEKENTKKDLLGIIKGMKVELSTVNV RTTKPPKRRPLKSLEATLGRLRRATEYAPKKRIEP LSPELVAAASAVADSLPFDKQTTKSELLSQLQQH EEESRAQRDAKRPKISFSNIISDMKVARSATARV RSRPELRIQFDEGYDNYPGQEKTDDLKKRKNIFT GKRLNIFDMMAVTKEAPETDTSPSLWDVEFAKQ LATVNEQPLQNGFEELIQWTKEGKLWEFPINNEA GFDDDGSEFHEHIFLEKHLESFPKQGPIRHFMELV TCGLSKNPYLSVKQKVEHIEWFRNYFNEKKDILK ESNIQFKLRPWKFLFRNN
3579	A	1725	445	RPRRGTHHFSCVLGSFRVSAMFPRVSTFLPLRP LSRHPLSSGSPETSAAAIMLLTVRHGTVRYRSSA LLARTKNNIQRYFGTNSVICSKKDKQSVRTEETS KETSESQDSEKENTKKDLLGIIKGMKVELSTVNV RTTKPPKRRPLKSLEATLGRLRRATEYAPKKRIEP LSPELVAAASAVADSLPFDKQTTKSELLSQLQQH EEESRAQRDAKRPKISFSNIISDMKVARSATARV RSRPELRIQFDEGYDNYPGQEKTDDLKKRKNIFT GKRLNIFDMMAVTKEAPETDTSPSLWDVEFAKQ LATVNEQPLQNGFEELIQWTKEGKLWEFPINNEA GFDDDGSEFHEHIFLEKHLESFPKQGPIRHFMELV TCGLSKNPYLSVKQKVEHIEWFRNYFNEKKDILK ESNIQFKLRPWKFLFRNN
3580	A	3673	1619	LYCVAPYSRHLLGRMSHLPMKLLRKKIEKRNLK LRQRNLKFQGASNLTLSETQNGDVSEETMGSRK VKKSKQKPMNVGLSETQNGGMSQEAVGNIKVT

SEQ ID NO:	Method	Predicted beginning nucleotide location corresponding to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion
				KSPQKSTVLTNGEAAMQSSNSESKKKKKKKK MVNDAEPDTKKAKTENKGKSEEESAETTKETEN NVEKPDNDEDESEVPSLPLGLTGAFEDTSFASLC NLVNENTLKAIKEMGFTNMTEIQHKSIRPLLEGR DLLAAAKTGSGKTLAFLIPAVELIVKLRFMPRNG TGVLILSPTRELAMQTFGVLKELMTHHVHTYGLI MGGSNRSAEAQKLGNGINIIVATPGRLLDHMQN TPGFMYKNLQCLVIDEADRILDVGFEEELKQIIKL LPTRRQTMLFSATQTRKVEDLARISLKKEPLYVG VDDDKANATVDGLEQGYVVCPSEKRFLLLFTFL KKNRKKKLMVFFSSCMSVKYHYELLNYIDLPVL AIHGKQKQNKRTTTFFQFCNADSGTLLCTDVAA RGLDIPEVDWIVQYDPPDDPKEYIHRVGRTARGL NGRGHALLILRPEELGFLRYLKQSKVPLSEFDFS WSKISDIQSQLEKLIEKNYFLHKSAQEAYKSYIRA YDSHSLKQIFNVNNLNLPQVALSFGFKVPPFVDL NVNSNEGKQKKRGGGGGGFGYQKTKKVEKSKIF KHISKKSSDSRQFSH
3581	A	23	453	LCRCICIKNITPHCLWDKVLSQFTYILDNLSNFMS HHPHSLRNSCLIRMDLLYWQFTIYTITFCFSHLSG RLTLSAQHISHRPCLLSYSLLFWKVHHLFLEGFPC SPRLDEMSFHQFPQHPVHVSVVHLPIVYKGSMT QVSPH
3582	A	3	950	TRGCGNKMAGKKNVLSSLAVYAEDSEPESDGEA GIEAVGSAAEEKGGLVSDAYGEDDFSRLGGDED GYEEEEDENSRQSEDDDSETEKPEADDPKDNTE AEKRDPQELVASFSERVRNMSPDEIKIPPEPPGRC SNHLQDKIQKLYERKIKEGMDMNYIIQRKKEFRN PSIYEKLIQFCAIDELGTNYPKDMFDPHGWSEDS YYEALAKAQKIEMDKLEKAKKERTKIEFVTGTK KGTTTNATSTTTTTASTAVADAQKRKSKWDSAI PVTTIAQPTILTTTATLPAVVTVTTSASGSKTTVIS AVGTIVKKAKQ
3583	A	3	950	TRGCGNKMAGKKNVLSSLAVYAEDSEPESDGEA GIEAVGSAAEEKGGLVSDAYGEDDFSRLGGDED GYEEEEDENSRQSEDDDSETEKPEADDPKDNTE AEKRDPQELVASFSERVRNMSPDEIKIPPEPPGRC SNHLQDKIQKLYERKIKEGMDMNYIIQRKKEFRN PSIYEKLIQFCAIDELGTNYPKDMFDPHGWSEDS YYEALAKAQKIEMDKLEKAKKERTKIEFVTGTK KGTTTNATSTTTTTASTAVADAQKRKSKWDSAI PVTTIAQPTILTTTATLPAVVTVTTSASGSKTTVIS AVGTIVKKAKQ
3584	A	3	1139	PGSTISSRADRLGAPVLAHPKMAERQEEQRGSPP LRAEGKADAEVKLILYHWTHSFSSQKVRLVIAE KALKCEEHDVSLPLSEHNEPWFMRLNSTGEVPV LIHGENIICEATQIIDYLEQTFLDERTPRLMPDKES MYYPRVQHYRELLDSLPMDAYTHGCILHPELTV DSMIPAYATTRIRSQIGNTESELKKLAEENPDLQE AYIAKQKRLKSKLLDHDNVKYLKKILDELEKVL DQVETELPRRNEETPEEGQQPWLCGESFTLADVS LAVTLHRLKFLGFARRNWGNGKRPNLETYYERV LKRKTFNKVLGHVNNILISAVLPTAFRVAKKRAP KVLGTTLVVGLLAGVGYFAFMLFRKRLGSMILA LRPRPNYF

SEQ ID NO:	Method	Predicted beginning nucleotide location corresponding to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methlonine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion
3585	A			RRHSPGSPAFAPSSRATAICPRAARAPATLLLALG AVLWPAAGAWELTILHTNDVHSRLEQTSEDSSK CVNASRCMGGVARLFTKVQQIRRAEPNVLLLDA GDQYQGTIWFTVYKGAEVAHFMNALRYDAMA LGNHEFDNGVEGLIEPLLKEAKFPILSANIKAKGP LASQISGLYLPYKVLPVGDEVVGIVGYTSKETPF LSNPGTNLVFEDEITALQPEVDKLKTLNVNKIIAL GHSGFEMDKLIAQKVRGVDVVVGGHSNTFLYT GNPPSKEVPAGKYPFIVTSDDGRKVPVVQAYAF GKYLGYLKIEFDERGNVISSHGNPILLNSSIPEDPS IKADINKWRIKLDNYSTQELGKTIVYLDGSSQSC RFRECNMGNLICDAMINNNLRHTDEMFWNHVS MCILNGGGIRSPIDERNNGTITWENLAAVLPFGG TFDLVQLKGSTLKKAFEHSVHRYGQSTGEFLQV GGIHVVYDLSRKPGDRVVKLDVLCTKCRVPSYD PLKMDEVYKVILPNFLANGGDGFQMIKDELLRH DSGDQDINVVSTYISKMKVIYPAVEGRIKFSTGS HCHGSFSLIFLSLWAVIFVLYQ
3586	A	1399	881	LSNKDVLSPQLKDENSKLRRKLNEVQSFSEAQTE MVRTLERKLEAKMIKEESDYHDLESVVQQVEQN LELMTKRAVKAENHVVKLKQEISLLQAQVSNFQ RENEALRCGQGASLTVVKQNADVALQNLRVVM NSAQASIEQLVSGAETLNLVAEILKSIDRISEVKD EEEDS
3587	A	88	1639	GCVGRGLPLPPRHPTPPSSSSSPFVLLAFLLLVRL DPAVSGKMAAPRPPPARLSGVMVPAPIQDLEAL RALTALFKEQRNRETAPRTIFQRVLDILKKSSHA VELACRDPSQVENLASSLQLITECFRCLRNACIEC SVNQNSIRNLDTIGVAVDLILLFRELRVEQESLLT AFRCGLQFLGNIASRNEDSQSIVWVHAFPELFLS CLNHPDKKIVAYSSMILFTSLNHERMKELEENLN IAIDVIDAYQKHPESEWPFLIITDLFLKSPELVQA MFPKLNNQERVTLLDLMIAKITSDEPLTKDDIPVF LRHAELIASTFVDQCKTVLKLASEEPPDDEEALA TIRLLDVLCEMTVNTELLGYLQVFPGLLERVIDL LRVIHVAGKETTNIFSNCGCVRAEGDISNVANGF KSHLIRLIGNLCYKNKDNQDKVNELDGIPLILDN CNISDSNPFLTQWVIYAIRNLTEDNSQNQDLIAK MEEQGLADASLLKKVGFEVEKKGEKLILKSTRD TPKP
3588	A		1462	DSPRNRFEILGRPTRTPTRPGPRPAMEDLDALLSD LETTTSHMPRSGAPKERPAEPLTPPPSYGHQPQT GSGESSGASGDKDHLYSTVCKPRSPKPAAPAAPP FSSSSGVLGTGLCELDRLLQELNATQFNITDEIMS QFPSSKVASGEQKEDQSEDKKRPSLPSSPSPGLPK ASATSATLELDRLMASLSDFRVQNHLPASGPTQP PVVSSTNEGSPSPPEPTGKGSLDTMLGLLQSDLSR RGVPTQAKGLCGSCNKPIAGQVVTALGRAWHPE HFVCGGCSTALGGSSFFEKDGAPFCPECYFERFSP RCGFCNQPIRHKMVTALGTHWHPEHFCCVSCGE PFGDEGFHEREGRPYCRRDFLQLFAPRCQGCQGP ILDNYISALSALWHPDCFVCRECFAPFSGGSFFEH EGRPLCENHFHARRGSLCATCGLPVTGRCVSAL GRRFHPDHFTCTFCLRPLTKGSFQERAGKPYCQP CFLKLFG

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SEQ ID NO:	Method	Predicted beginning	Predicted end nucleotide	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine,
110.		nucleotide	location	I=Isoleucine, K=Lysine, L=Leucine, M=Methionine,
		location	corresponding	N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine,
	1	corresponding	to last amino	T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine,
		to first amino acid residue of	acid residue of	X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion
		peptide	peptide sequence	/-bossing undeading uper non
	1	sequence	100420000	
3589	A	226	6793	SPPKKSRKCNLSFRLISAERWRFFLLILMEMPRKP
	İ	Į		RLTLFVQRRIENIATEREFDPEEFYYLLEAAEGHA
		1		KEGQGIKTDIPRYIISQLGLNKDPLEEMAHLGNY
			•	DSGTAETPETDESVSSSNASLKLRRKPRESDFETI
	1	}		KLISNGAYGAVYFVRHKESRQRFAMKKINKQNL
				ILRNQIQQAFVERDILTFAENPFVVSMYCSFETRR
				HLCMVMEYVEGGDCATLMKNMGPLPVDMARM
	İ			YFAETVLALEYLHNYGIVHRDLKPDNLLVTSMG
				HIKLTDFGLSKVGLMSMTTNLYEGHIEKDAREFL
	}	ļ	ļ	DKQVCGTPEYIAPEVILRQGYGKPVDWWAMGII
				LYEFLVGCVPFFGDTPEELFGQVISDEINWPEKDE
l			}	APPPDAQDLITLLLRQNPLERLGTGGAYEVKQHR
l				FFRSLDWNSLLRQKAEFIPQLESEDDTSYFDTRSE
				KYHHMETEEEDDTNDEDFNVEIRQFSSCSHRFSK
	1			VFSSIDRITQNSAEEKEDSVDKTKSTTLPSTETLS
			Į	WSSEYSEMQQLSTSNSSDTESNRHKLSSGLLPKL
	į			AISTEGEQDEAASCPGDPHEEPGKPALPPEECAQ
				EEPEVTTPASTISSSTLSVGSFSEHLDQINGRSECV DSTDNSSKPSSEPASHMARQRLESTEKKKISGKV
				TKSLSASALSLMIPGDMFAVSPLGSPMSPHSLSSD
i				PSSSRDSSPSRDSSAASASPHQPIVIHSSGKNYGFT
				IRAIRVYVGDSDIYTVHHIVWNVEEGSPACQAGL
}				KAGDLITHINGEPVHGLVHTEVIELLLKSGNKVSI
				TTTPFENTSIKTGPARRNSYKSRMVRRSKKSKKK
				ESLERRRSLFKKLAKQPSPLLHTSRSFSCLNRSLS
	l			SGESLPGSPTHSLSPRSPTPSYRSTPDFPSGTNSSO
				SSSPSSSAPNSPAGSGHIRPSTLHGLAPKLGGQRY
ļ				RSGRRKSAGNIPLSPLARTPSPTPQPTSPQRSPSPL
ĺ		†		LGHSLGNSKIAQAFPSKMHSPPTIVRHIVRPKSAE
	ļ			PPRSPLLKRVQSEEKLSPSYGSDKKHLCSRKHSL
ļ				EVTQEEVQREQSQREAPLQSLDENVCDVPPLSRA
		1		RPVEQGCLKRPVSRKVGRQESVDDLDRDKLKAK
				VVVKKADGFPEKQESHQKFHGPGSDLENFALFK
1		1		LEEREKKVYPKAVERSSTFENKASMQEAPPLGSL
]			LKDALHKQASVRASEGAMSDGPVPAEHRQGGG
				DFRRAPAPGTLQDGLCHSLDRGISGKGEGTEKSS
				QAKELLRCEKLDSKLANIDYLRKKMSLEDKEDN
	1	}		LCPVLKPKMTAGSHECLPGNPVRPTGGQQEPPPA
				SESRAFVSSTHAAQMSAVSFVPLKALTGRVDSGT
]				EKPGLVAPESPVRKSPSEYKLEGRSVSCLEPIEGT
1				LDIALLSGPQASKTELPSPESAQSPSPSGDVRASV
1			,	PPVLPSSSGKKNDTTSARELSPSSLKMNKSYLLEP
				WFLPPSRGLQNSPAVSLPDPEFKRDRKGPHPTAR
				SPGTVMESNPQQREGSSPKHQDHTTDPKLLTCLG
1				QNLHSPDLARPRCPLPPEASPSREKPGLRESSERG PPTARSERSAARADTCREPSMELCFPETAKTSDN
				SKNLLSVGRTHPDFYTQTQAMEKAWAPGGKTN
				HKDGPGEARPPPRDNSSLHSAGIPCEKELGKVRR
1	}			GVEPKPEALLARRSLQPPGIESEKSEKLSSFPSLQ
		[KDGAKEPERKEQPLQRHPSSIPPPPLTAKDLSSPA
				ARQHCSSPSHASGREPGAKPSTAEPSSSPQDPPKP
}]			VAAHSESSSHKPRPGPDPGPPKTKHPDRSLSSQK
				PSVGATKGKEPATQSLGGSSREGKGHSKSGPDVF
]				PATPGSQNKASDGIGQGEGGPSVPLHTDRAPLDA
		}		KPQPTSGGRPLEVLEKPVHLPRPGHPGPSEPADQ
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SEQ ID NO:	Method	Predicted beginning	Predicted end nucleotide	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine,
110.		nucleotide location corresponding to first amino acid residue of peptide	location corresponding to last amino acid residue of peptide sequence	I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion
		sequence		KLSAVGEKQTLSPKHPKPSTVKDCPTLCKQTDN RQTDKSPSQPAANTDRRAEGKKCTEALYAPAEG DKLEAGLSFVHSENRLKGAERPAAGVGKGFPEA RGKGPGPQKPPTEADKPNGMKRSPSATGQSSFRS TALPEKSLSCSSSFPETRAGVREASAASSDTSSAK AAGGMLELPAPSNRDHRKAQPAGEGRTHMTKS DSLPSFRVSTLPLESHHPDPNTMGGASHRDRALS VTATVGETKGKDPAPAQPPPARKQNVGRDVTKP SPAPNTDRPISLSNEKDFVVRQRRGKESLRSSPHK KAL
3590	A	3	935	RATTRPKNEVQDYVSVEYLSPHMGGTDPFKYSY PPLVDDDFQTPLCENGPITSEDETSSKEDIESDGK ETLETISNEEQTPLLKKINPTESTSKAEENEKVDS KVKAFKKPLSVFKGPLLHISPAEELYFGSTESGEK KTLIVLTNVTKNIVAFKVRTTAPEKYRVKPSNSS CDPGASVDIVVSPHGGLTVSAQDRFLIMAAEME QSSGTGPAELTQFWKEVPRNKVMEHRLRCHTVE SSKPNTLTLKDNAFNMSDKTSEDICLQLSRLLES NRKLEDQVQRCIWFQQLLLSLTMLLLAFVTSFFY LLYS
3591	A	303	2	GGSWGPLCPVSPAMSLSDPGLGYHPTCWTLRWP PLCSLHALHVFHCLFSSRLGTPVSPRLAMDPNCS CEAGGSCACAGSCKCKKCKCTSCKKSCCSCCPL
3592	A .	1052	1779	GKTMMRKMLLAAALSVTAMTAHADYQCSVTP RDDVIVSPQTVQVKGENGNLVITPDGNVMYNGK QYSLNAAQREQAKDYQAELRSTLPWIDEGAKSR VEKARIALDKIIVQEMGESSKMRSRLTKLDAQVK EQMNRIIETRSDGLTFHYKAIDQVRAEGQQLVNQ AMGGILQDSINEMGAKAVLKSGGNPLQNVLGSL GGLQSSIQTEWKKQEKDFQQFGKDVCSRVVTLE DSRKALVGNLK
3593		3	1837	LSFEKVDIQTDNDLTKEMYEGKENVSFELQRDFS QETDFSEASLLEKQQEVHSAGNIKKEKSNTIDGT VKDETSPVEECFFSQSSNSYQCHTITGEQPSGCTG LGKSISFDTKLVKHEIINSEERPFKCEELVEPFRCD SQLIQHQENNTEEKPYQCSECGKAFSINEKLIWH QRLHSGEKPFKCVECGKSFSYSSHYITHQTIHSGE KPYQCKMCGKAFSVNGSLSRHQRIHTGEKPYQC KECGNGFSCSSAYITHQRVHTGEKPYECNDCGK AFNGNAKLIQHQRIHTGEKPYECNECGKGFRCSS QLRQHQSIHTGEKPYQCKECGKGFNNNTKLIQH QRIHTASLAEQLFKASGNHPNWGCCLTISSPGPS VYGPKMNMRGAPNSRLAGGREKRTQDTDFGQC SFLPSHSPSCFEPWNVTDYDSSWYRQKQVLSGV WSSPLSILKLPRTLIRISIHIQEMDTPGEMLMTGR GSLGPTLTTEAPAAAQPGKQGPPGTGRCLQAPGT EPGEQTPEGARELSPLQESSSPGGVKAEEEQRAG AEPGTRPSLARSDDNDHEVGALGLQQGKSPGAG NPEPEQDCAARAPVRAEAVRRMPPGAEAGSVVL DD
3594	A	39	261	RAAMMDTSRVQPIKLAIVIKVLGRTGSQGQCTQ VRVEFMDDTSRSIIRSVKGPVREGDVLTLLESERE ARRLR
3595	A	973	68	GRVGTKHQMADDAGAAGGPGGPGGPGMGNRG GFRGGFGSGIRGRGRGRGRGRGRGRARGGKAE

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SEQ ID NO:	Method	Predicted beginning nucleotide location corresponding to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion
				DKEWMPVTKLGRLVKDMKIKSLEEIYLFSLPIKE SEIIDFFLGASLKDEVLKIMPVQKQTRAGQRTRF KAFVAIGDYNGHVGLGVKCSKEVATAIRGAIILA KLSIVPVRRGYWGNKIGKPHTVPCKVTGRCGSV LVRLIPAPRGTGIVSAPVPKKLLMMAGIDDCYTS ARGCTATLGNFAKATFDAISKTYSYLTPDLWKE TVFTKSPYQEFTDHLVKTHTRVSVQRTQAPAVA TT
3596		106	2960	DERRVGAADMFGRSRSWVGGGHGKTSRNIHSL DHLKYLYHVLTKNTTVTEQNRNLLVETIRSITEIL IWGDQNDSSVFDFFLEKNMFVFFLNILRQKSGRY VCVQLLQTLNILFENISHETSLYYLLSNNYVNSII VHKFDFSDEEIMAYYISFLKTLSLKLNNHTVHFF YNEHTNDFALYTEAIKFFNHPESMVRIAVRTITL NVYKVSLDNQAMLHYIRDKTAVPYFSNLVWFIG SHVIELDDCVQTDEEHRNRGKLSDLVAEHLDHL HYLNDILIINCEFLNDVLTDHLLNRLFLPLYVYSL ENQDKGGERPKISLPVSLYLLSQVFLIIHHAPLVN SLAEVILNGDLSEMYAKTEQDIQRSSAKPSIRCFI KPTETLERSLEMNKHKGKRRVQKRPNYKNVGEE EDEEKGPTEDAQEDAEKAKGTEGGSKGIKTSGES EEIEMVIMERSKLSELAASTSVQEQNTTDEEKSA AATCSESTQWSRPFLDMVYHALDSPDDDYHALF VLCLLYAMSHNKGMDPEKLERIQLPVPNAAEKT TYNHPLAERLIRIMNNAAQPDGKIRLATLELSCL LLKQQVLMSAGCIMKDVHLACLEGAREESVHLV RHFYKGEDIFLDMFEDEYRSMTMKPMNVEYLM MDASIILPPTGTPLTGIDFVKRLPCGDVEKTRRAI
				MDASILLPPTGTPLTGIDFVKRLPCGDVEKTRRAI RVFFMLRSLSLQLRGEPETQLPLTREEDLIKTDDV LDLNNSDLIACTVITKDGGMVQRSLAVDIYQMS LVEPDVSRLGWGVVKFAGLLQDMQVTGVEDDS RALNITIHKPASSPHSKPFPILQATFIFSDHIRCIIAK QRLAKGRIQARRMKMQRIAALLDLPIQPTTEVLG FGLGSSTSTQHLPFRFYDQGRRGSSDPTVQRSVF ASVDKVPGFAVAQCINEHSSPSLSSQSPPSASGSP SGSGSTSHCDSGGTSSSSTPSTAQSPAGIGHVTQ
3597	A	427	277	GVRRIQHHWAQMHECNVHTYASLFCLFLLHTG KLCCLNSHRHFHCIKYSK
3598	A	1	503	FRPRTKKATAMYLEHYLDSIENLPCELQRNFQL MRELDQRTEDKKAEIDILAAEYISTVKTLSPDQR VERLQKIQNAYSKCKEYSDDKVQLAMQTYEMV DKHIRRLDADLARFEADLKDKMEGSDFESSGGR GLKKGRGQKEKRGSRGRGRRTSEEDTPKKKKH KGG
3599	A		3907	KTITALAFSPDGKYLVTGESGHMPAVRVWDVAE HSQVAELQEHKYGVACVAFSPSAKYIVSVGYQH DMIVNVWAWKKNIVVASNKVSSRVTAVSFSED CSYFVTAGNRHIKFWYLDDSKTSKVNATVPLLG RSGLLGELRNNLFTDVACGRGKKADSTFCITSSG LLCEFSDRRLLDKWVELRVYPEVKDSNQACLPP SSFITCSSDNTIRLWNTESSGVHGSTLHRNILSSDL IKIIYVDGNTQALLDTELPGGDKADASLLDPRVGI RSVCVSPNGQHLASGDRMGTLRVHELQSLSEML KVEAHDSEILCLEYSKPDTGLKLLASASRDRLIH VLDAGREYSLQQTLDEHSSSITAVKFAASDGQVR

SEQ ID NO:	Method	Predicted beginning nucleotide location corresponding to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, I=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion
				MISCGADKSIYFRTAQKSGDGVQFTRTHHVVRK TTLYDMDVEPSWKYTAIGCQDRNIRIFNISSGKQ KKLFKGSQGEDGTLIKVQTDPSGIYIATSCSDKNL SIFDFSSGECVATMFGHSEIVTGMKFSNDCKHLIS VSGDSCIFVWRLSSEMTISMRQRLAELRQRQRGG KQQGPSSPQRASGPNRHQAPSMLSPGPALSSDSD KEGEDEGTEEELPALPVLAKSTKKALASVPSPAL PRSLSHWEMSRAQESVGFLDPAPAANPGPRRRG RWVQPGVELSVRSMLDLRQLETLAPSLQDPSQD SLAIIPSGPRKHGQEALETSLTSQNEKPPRPQASQ PCSYPHIRLLSQEEGVFAQDLEPAPIEDGIVYPEP SDNPTMDTSEFQVQAPARGTLGRVYPGSRSSEK HSPDSACSVDYSSSCLSSPEHPTEDSESTEPLSVD GISSDLEEPAEGDEEEEEEEGGMGPYGLQEGSPQ TPDQEQFLKQHFETLASGAAPGAPVQVPERSESR SISSRFLLQVQTRPLREPSPSSSSLALMSRPAQVPQ ASGEQPRGNGANPPGAPPEVEPSSGNPSPQQAAS VLLPRCRLNPDSSWAPKRVATASPFSGLQKAQS VHSLVPQERHEASLQAPSPGALLSREIEAQDGLG SLPPADGRPSRPHSYQNPTTSSMAKISRSISVGEN LGLVAEPQAHAPIRVSPLSKLALPSRAHLVLDIPK PLPDRPTLAAFSPVTKGRAPGEAEKPGFPVGLGK AHSTTERWACLGEGTTPKPRTECQAHPGPSSPCA QQLPVSSLFQGPENLQPPPPEKTPNPMECTKPGA ALSQDSEPAVSLEQCEQLVAELRGSVRQAVRLY HSVAGCKMPSAEQSRIAQLLRDTFSSVRQELEAV AGAVLSSPGSSPGAVGAEQTQALLEQYSELLLRA VERRMERKL
3600	A	1688	916	IPGSTISCSMALCEAAGCGSALLWPRLLLFGDSIT QFSFQQGGWGASLADRLVRKCDVLNRGFSGYN TRWAKIILPRLIRKGNSLDIPVAVTIFFGANDSAL KDENPKQHIPLEEYAANLKSMVQYLKSVDIPENR VILITPTPLCETAWEEQCIIQGCKLNRLNSVVGEY ANACLQVAQDCGTDVLDLWTLMQDSQDFSSYL SDGLHLSPKGNEFLFSHLWPLIEKKVSSLPLLLPY WRDVAEAKPELSLLGDGDH
3601	A	44	223	VHFPLIPQLAKCFWTMNRAARNKSEKRYYSEFL QIAHLFNYGLSSFLREFIIFLIKLLQ
3602	A	37	1124	VPKPASGKRRLEFRPQDSKACAATPHSPGRITSR TRGSQKVRSVPPRLPWAQASASTDWEGLRGVPG PALRENFLEAAASGRSGRTPTGGVGFRDVGGP HFPIFPAAHFLWCNLHTPRRPACNAPWHSPVGEI SPPPRESQLRRDPEVHFESPAHPLGFRLLPGRGLP ANAVTVETAAMAAPRQIPSHIVRLKPSCSTDSSF TRTPVPTVSLASRELPVSSWQVTEPSSKNLWEQI CKEYEAEQPPFPEGYKVKQEPVITVAPVEEMLFH GFSAEHYFPVSHFTMISRTPCPQDKSETINPKTCS PKEYLETFIFPVLLPGMASLLHQAKKEKCFEVVL QMTPSGGKACVWGHLPSSSHTI
3603	A	286	587	NISNKAEVSSHPSVISHSMDSFGQPRPEDNQSVLR RMQKKYWKTKQVFIKATGKKEDEHLVASDAEL DAKLEVFHSVQETCTELLKIIEKYQLRLNGMKS
3604	A	103	2440	QPRRRVFPAAGRGPGRKCSQWGRQASVSFEDVT VDFSKEEWQHLDPAQRRLYWDVTLENYSHLLS VGYQIPKSEAAFKLEQGEGPWMLEGEAPHQSCS

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SEQ ID NO:	Method	Predicted beginning nucleotide location corresponding to first amino acid residue of peptide	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion
		sequence		GEAIGKMQQQGIPGGIFFHCERFDQPIGEDSLCSI LEELWQDNDQLEQRQENQNNLLSHVKVLIKERG YEHKNIEKIIHVTTKLVPSIKRLHNCDTILKHTLN SHNHNRNSATKNLGKIFGNGNNFPHSPSSTKNEN AKTGANSCEHDHYEKHLSHKQAPTHHQKIHPEE KLYVCTECVMGFTQKSHLFEHQRIHAGEKSREC DKSNKVFPQKPQVDVHPSVYTGEKPYLCTQCGK VFTLKSNLITHQKIHTGQKPYKCSECGKAFFQRS DLFRHLRIHTGEKPYECSECGKGFSQNSDLSIHQ KTHTGEKHYECNECGKAFTRKSALRMHQRIHTG EKPYVCADCGKAFIQKSHFNTHQRIHTGEKPYEC SDCGKSFTKKSQLHVHQRIHTGEKPYICTECGKV FTHRTNLTTHQKTHTGEKPYMCAECGKAFTDQS NLIKHQKTHTGEKPYKCNGCGKAFIWKSRLKIH QKSHIGERHYECKDCGKAFIQKSTLSVHQRIHTG EKPYVCPECGKAFIQKSHFIAHHRIHTGEKPYECS DCGKCFTKKSQLRVHQKIHTGEKPNICAECGKAF TDRSNLITHQKIHTREKPYECGDCGKTFTWKSRL NIHQKSHTGERHYECSKCGKAFIQKATLSMHQII HTGKKPYACTECQKAFTDRSNLIKHQKMHSGEK RYKASD
3605	A	3	322	SFRMSGRGKGGKGLGKGGAKRHRKVLRDNIQGI TKPAIRRLARRGGVKRISGLIYEETRGVLKVFLEN VIRDAVTYTEHAKRKTVTAMDVVYALKRQGRT LYGFGG
3606	A		1749	VPVTAEAKLMGFTQGCVTFEDVAIYFSQEEWGL LDEAQRLLYRDVMLENFALITALVCWHGMEDE ETPEQSVSVEGVPQVRTPEASPSTQKIQSCDMCV PFLTDILHLTDLPGQELYLTGACAVFHQDQKHHS AEKPLESDMDKASFVQCCLFHESGMPFTSSEVG KDFLAPLGILQPQAIANYEKPNKISKCEEAFHVGI SHYKWSQCRRESSHKHTFFHPRVCTGKRLYESS KCGKACCCECSLVQLQRVHPGERPYECSECGKS FSQTSHLNDHRRIHTGERPYVCGQCGKSFSQRAT LIKHHRVHTGERPYECGECGKSFSQSSNLIEHCRI HTGERPYECDECGKAFGSKSTLVRHQRTHTGEK PYECGECGKLFRQSFSLVVHQRIHTTARPYECGQ CGKSFSLKCGLIQHQLIHSGARPFECDECGKSFSQ RTTLNKHHKVHTAERPYVCGECGKAFMFKSKL VRHQRTHTGERPFECSECGKFFRQSYTLVEHQKI HTGLRPYDCGQCGKSFIQKSSLIQHQVVHTGERP YECGKCGKSFTQHSGLILHRKSHTVERPRDSSKC GKPYSPRSNIV
3607	A	92	331	AMAGPGPGPGDPDEQYDFLFKLVLVGDASVGKT CVVQRFKTGAFSERQGSTIGVDFTMKTLEIQGKR VKLQIWDTAGQER
3608	A	545	379	AIKGYIHLSAPRNRYMHTTASNGRMLFMKVTM YMRRGVQIMGWSVRMAFMACFTQ
3609	A	118	873	VWMAWQVSLLELEDRLQCPICLEVFKESLMLQC GHSYCKGCLVSLSYHLDTKVRCPMCWQVVDGS SSLPNVSLAWVIEALRLPGDPEPKVCVHHRNPLS LFCEKDQELICGLCGLLGSHQHHPVTPVSTVCSR MKEELAALFSELKQEQKKVDELIAKLVKNRTRIV NESDVFSWVIRREFQELRHPVDEEKARCLEGIGG HTRGLVASLDMQLEQAQGTRERLAQAECVLEQF

SEQ ID NO:	Method	Predicted beginning nucleotide location corresponding to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \text{\tex{\tex
				GNEDHHEFIWKFHSMASR
3610	Α	2	987	DPRVRPPLLQPPPPLLPRLVILKMAPLDLDKYVEI
				ARLCKYLPENDLKRLCDYVCDLLLEESNVQPVS TPVTVCGDIHGQFYDLCELFRTGGQVPDTNYIFM GDFVDRGYYSLETFTYLLALKAKWPDRITLLRG NHESRQITQVYGFYDECQTKYGNANAWRYCTK VFDMLTVAALIDEQILCVHGGLSPDIKTLDQIRTI ERNQEIPHKGAFCDLVWSDPEDVDTWAISPRGA GWLFGAKVTNEFVHINNLKLICRAHQLVHEGYK
ł			i	FMFDEKLVTVWSAPNYCYRCGNIASIMVFKDVN
3611	A	2459	869	TREPKLFRAVPDSERVIPPRTTTPYFL AEKMTAELREAMALAPWGPVKVKKEEEEEENF PGQASSQQVHSENIKVWAPVQGLQTGLDGSEEE EKGQNISWDMAVVLKATQEAPAASTLGSYSLPG TLAKSEILETHGTMNFLGAETKNLQLLVPKTEIC EEAEKPLIISERIQKADPQGPELGEACEKGNMLK RQRIKREKKDFRQVIVNDCHLPESFKEEENQKCK KSGGKYSLNSGAVKNPKTQLGQKPFTCSVCGKG FSQSANLVVHQRIHTGEKPFECHECGKAFIQSAN LVVHQRIHTGQKPYVCSKCGKAFTQSSNLTVHQ
3612	A	318	2245	KIHSLEKTFKCNECEKAFSYSSQLARHQKVHITE KCYECNECGKTFTRSSNLIVHQRIHTGEKPFACN DCGKAFTQSANLIVHQRSHTGEKPYECKECGKA FSCFSHLIVHQRIHTAEKPYDCSECGKAFSQLSCL IVHQRIHSGDLPYVCNECGKAFTCSSYLLIHQRIH NGEKPYTCNECGKAFRQRSSLTVHQRTHTGEKP YECEKCGAAFISNSHLMRHHRTHLVE SPMAEAALVNTPQIPMVTEEFVKPSQGHVTFEDI
				AVYFSQEEWGLLDEAQRCLYHDVMLENFSLMA SVGCLHGIEAEEAPSEQTLSAQGVSQARTPKLGP SIPNAHSCEMCILVMKDILYLSEHQGTLPWQKPY TSVASGKWFSFGSNLQQHQNQDSGEKHIRKEESS ALLLNSCKIPLSDNLFPCKDVEKDFPTILGLLQHQ TTHSRQEYAHRSRETFQQRRYKCEQVFNEKVHV TEHQRVHTGEKAYKRREYGKSLNSKYLFVEHQR THNAEKPYVCNICGKSFLHKQTLVGHQQRIHTRE RSYVCIECGKSLSSKYSLVEHQRTHNGEKPYVCN VCGKSFRHKQTFVGHQQRIHTGERPYVCMECGK SFIHSYDRIRHQRVHTGEGAYQCSECGKSFIYKQ SLLDHHRIHTGERPYECKECGKAFIHKKRLLEHQ RIHTGEKPYVCIICGKSFIRSSDYMRHQRIHTGER AYECSDCGKAFISKQTLLKHHKIHTRERPYECSE CGKGFYLEVKLLQHQRIHTREQLCECNECGKVF SHQKRLLEHQKVHTGEKPCECSECGKCFRHRTS LIQHQKVHSGERPYNCTACEKAFIYKNKLVEHQ RIHTGEKPYECGKCGKAFNKRYSLVRHQKVHIT EEP
3613	A	817	3345	NQSHPDSETVTVEGGRRKMKSNQERSNECLPPK KREIPATSRSSEEKAPTLPSDNHRVEGTAWLPGN PGGRGHGGGRHGPAGTSVELGLQQGIGLHKALS TGLDYSPPSAPRSVPVATTLPAAYATPQPGTPVSP VQYAHLPHTFQFIGSSQYSGTYASFIPSQLIPPTAN PVTSAVASAAGATTPSQRSQLEAYSTLLANMGS LSQTPGHKAEQQQQQQQQQQQQQQQQQQQQQQQQQQQQQQQQQQQQ

SEQ ID NO:	Method	Predicted beginning nucleotide location corresponding to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \text{\tex{\tex
		sequence		PAQQNQYVHISSSPQNTGRTASPPAIPVHLHPHQ TMIPHTLTLGPPSQVVMQYADSGSHFVPREATK KAESSRLQQAIQAKEVLNGEMEKSRRYGAPSSA DLGLGKAGGKSVPHPYESRHVVVHPSPSDYSSR DPSGVRASVMVLPNSNTPAADLEVQQATHREAS PSTLNDKSGLHLGKPGHRSYALSPHTVIQTTHSA SEPLPVGLPATAFYAGTQPPVIGYLSGQQQAITY AGSLPQHLVIPGTQPLLIPVGSTDMEASGAAPAIV TSSPQFAAVPHTFVTTALPKSENFNPEALVTQAA YPAMVQAQIHLPVVQSVASPAAAPPTLPPYFMK GSIIQLANGELKKVEDLKTEDFIQSAEISNDLKIDS STVERIEDSHSPGVAVIQFAVGEHRAQVSVEVLV EYPFFVFGQGWSSCCPERTSQLFDLPCSKLSVGD VCISLTLKNLKNGSVKKGQPVDPASVLLKHSKA DGLAGSRHRYAEQENGINQGSAQMLSENGELKF PEKMGLSAAPFLTKIEPSKPAATRKRRWSAPESR KLEKSEDEPPLTLPKPSLIPQEVKICIEGRSNVGK
3614	A	3	114	FFESRLRCKCCEPRGSWARFGCWRLQPEFKPKQ LEG
3615	A	3	1603	DAWALTNQFSDSKQHIEVLKESLTAKEQRAAILQ TEVDALRLRLEEKETMLNKKTKQIQDMAEEKGT QAGEIHDLKDMLDVKERKVNVLQKKIENLQEQL RDKEKQMSSLKERVKSLQADTTNTDTALTTLEE ALAEKERTIERLKEQRDRDEREKQEEIDNYKKDL KDLKEKVSLLQGDLSEKEASLLDLKEHASSLASS GLKKDSRLKTLEIALEQKKEECLKMESQLKKAH EAALEARASPEMSDRIQHLEREITRYKDESSKAQ AEVDRLLEILKEVENEKNDKDKKIAELESLTSRQ VKDQNKKVANLKHKEQVEKKKSAQMLEEARRR EDNLNDSSQQLQDSLRKKDDRIEELEEALRESVQ ITAEREMVLAQEESARTNAEKQVEELLMAMEKV KQELESMKAKLSSTQQSLAEKETHLTNLRAERR KHLEEVLEMKQEALLAAISEKDANIALLELSSSK KKTQEEVAALKREKDRLVQQLKQQTQNRMKLM ADNYEDDHFKSSHSNQTNHKPSPDQDEEEGIWA
3616		244	1420	RRRWRARGGLVPTLAWAEATGAYVPGRDKPDL PTWKRNFRSALNRKEGLRLAEDRSKDPHDPHKI YEFVNSGVGDFSQPDTSPDTNGGGSTSDTQEDIL DELLGNMVLAPLPDPGPPSLAVAPEPCPQPLRSPS LDNPTPFPNLGPSENPLKRLLVPGEEWEFEVTAF YRGRQVFQQTISCPEGLRLVGSEVGDRTLPGWP VTLPDPGMSLTDRGVMSYVRHVLSCLGGGLAL WRAGQWLWAQRLGHCHTYWAVSEELLPNSGH GPDGEVPKDKEGGVFDLGPFIVGSLGPPDLITFTE GSGRSPRYALWFCVGESWPQDQPWTKRLVMVK VVPTCLRALVEMARVGGASSLENTVDLHISNSHP LSLTSDQYKAYLQDLVEGMDFQGPGES
3617	Α	852	304	RGGLLSKMARVLKAAAANAVGLFSRLQAPIPTV RASSTSQPLDQVTGSVWNLGRLNHVAIAVPDLE KAAAFYKNILGAQVSEAVPLPEHGVSVVFVNLG NTKMELLHPLGRDSPIAGFLQKNKAGGMHHICIE VDNINAAVMDLKKKKIRSLSEEVKIGAHGKPVIF LHPKDCGGVLVELEQA
3618	A	3	5992	DNIDETYGVNVQFESDEEEGDEDVYGEVREEAS DDDMEGDEAVVRCTLSANMYVDEILVWCASEL

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SEQ ID NO:	Method	Predicted beginning nucleotide location corresponding to first amino acid residue of peptide	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \text{\tex{\tex
		sequence		
		acid residue of	peptide	
				WLSPALAAMELAQMVTQAMWSEDSYLRRLPPF PSGLFKRCTDKGVESVFDIMEMEDEERNALLQLT DSQIADVARFCNRYPNIELSYEVVDKDSIRSGGP VVVLVQLEREEEVTGPVIAPLFPQKREEGWWVV

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NO: Methodic Succession	SEQ ID	Method	Predicted	Predicted end	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid,
IGDAKSNSLISIKRLTLQQKAKVKLDFVAPATGG RHNTLYFMSDAYMGCDQEYKFSVDVKRAETDS DSD DND DND DND DND DND DND D	_	Wieinod	beginning nucleotide location corresponding to first amino acid residue of	nucleotide location corresponding to last amino acid residue of peptide	E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion,
RHNTLYFMSDAYMGCDQEYKFSVDVKEAETDS DSD 3619 A 3 5992 DNIDETYGVNVQFSDEEEGDEDVYGEVREAS DDDMAGDBAVVRCTLSANMYVDEIL.WCASEL NIPEFFLESPHKKVGYGLSSRTVLQGGGKVUS GRPLLVASGELMSSKKKDLHPRDIDAFWLQRQL VELLGHTFEDELVAKADEVLEILKTASDRECENQL VELLGHTTEDFILVARQHRMMILYCTLLASAQSE AEKERIMGKMEADPELSKFLYQLHETEKEDLIRE ERSRERRYRQSRMDTDLETMDLDQGGEALAPRQ VLDLEDLVFTQGSHFMANKRCQLDOSFRRQRK GYEEVHPALKPFGSEGULPVEKLFKYAQA GFEGFKTLNRIQSKLYRAALETDENLLLCAPTGA GKTNVALMCMLREIGKHINMDGTINVDDFKIIYI APMRSLVQEMVGSFGKRLATYGTIVAELTGDHQ LCKEEISATQIIVCTPEKWDITRKGGERTYTQL.V RLILLDEHILLHDDRGPVLEALVARARINNEMTQE DVRLIGLSAATLPNYEDVATELRVDPAKGLFYFDN SRPPVPLEQTYVGUTEKKAKIRFGJMMEIVVEKIM EHAGKNQVLVFVHSRKETGKTARAIRDMCLEKD TLGLFLREGSASTEVLRTEARQCKNLELKDLLPY GRÄHHAGMTRVDRTLVEDLFGDKHQVLVSTA TLAWGVNLPAHTVIIKGTGYSPEKGRWTELGA LDLQMLGRAGRPQVTDKGEGLITTSIGGLQVYL SLLNQQLPIESQMYSKLPDMLNAEIVLGNVQNA KDANNULGYANLYRMLRSPTLYGSHDDLKG DPLLDQRRLDLVHTAALMLDKNNLKYDKKTON FQVTELGRLASHYYITNDTVQTYNQLLKPTLSEIE LFVYFSLSSEFKNITVREEKKLELGKLLERPY ESIEEPSAKINVLQAFISQLKLEGFALMADMYY VTQSAGRLMRAFEVLNGWAQLTDKITLNCK MIDKRAMYOSMCPLRQFRKLPEEVVKKIEKKONFP FERLYDLNHNEIGELIRMPKMGKTHKYVLFTK ESIEEPSAKINVLQAFISQLKLEGFALMADMYY VTQSAGRLMRAFEVLNGWAQLTDKITLNCK MIDKRAMYOSMCPLRQFRKLPEEVVKKIEKKONFP FERLYDLNHNEIGELIRMPKMGKTHKYVLLFY ELSVHLQPITRSTLKVELTTIPDFQWDEKVHGSS EAFWILVEDVDSEVILHHEYFLLKAKYAQDEHLI TFFVYPFEPLPPQYTFRVSDRWLSCETQLCYPY MDWYSKFQDRLINKKVVLLTGETSTDLKLLGKG NIISTPEKWDLSRSRWKQKKNVONNINLEVVUDEV HLLGGENGPVLEVICSRMKYVSCGTQLCYPY MDWYSKFQDRLINKKVVLLTGETSTDLKLLGKG NIISTPEKWDLSRSRWKQKKNVONNINLEVVUDEV HLLGGENGPVLEVICSRMKYNSSQIERPIRIVALSSS LSNAKGCSATSTFNPHRVNPVPLELH QGFNISHTOTIRLLSMAKPYFHAITKHSPKKPVTVP VPSRKQTRLTADILTTCAADIQRQRFLHCTEKDL PYYLEKLSDSTLKETLLINGVGYLHEGLSPMERRIL VEQLESSGAJQVVVASRSLCWGMNVAAHLVIIM DTLYYMGKHHAYVDRYTDVLQMGHANREIL DFYLEKLSDSTLKETLLINGVGYLHEGLSPMERRIL VEQLESSGAJQVVVASRSLCWGMNVAAHLVIIM DTLYYMGKHHAYVDRYDYDVLQMGHANREILD HCMHIDHNABIVTKTENKODAADVILTWITTELL FFMSLNAMKTKWRGLEIISNAAEZPWIPITHHHEDL FFMSLNAMKTKWRGLEIISNAAEZPWIPITHHEDL FFMSLNAMKTKWRGLEIISNAAEZPWIPITHHEDL FFMSLNAMKTKWRGLEIISNAAEZPWIPITHHEDL FFMSLNA					·
3 5992 DNIDETYGVNYQFESDEEGDEDVYGVYREAS DDDMEGDBAVVRCTLSANMYVDEILVWCASEL NIPEFFILESPHKKVGYGLSSRTWLQGGGKVIEA GRDLLVASGELMSSKKKDLHPRDIDAFWLQRQL SRRYDDAIVOSGKADEVLEILKTASDDRECENQL VLLLGFNTFDFIKVLRQHRMMILYCTLLASAQSE AEKERIMGKMBADPELSKHLYQLHETEKBLIRE ERSRRERVRQSRMDTDLETMDLDQGGEALAPRQ VLDLEDLVFTQGSHFMANKRCQLPDGSFRRQRK GYEEVHYPALKPRPFOSEEGLLPVEKLPKYAQA GFEGFKTLNRIQSKLYRAALETDENLLLCAPTGA GKTNVALMCMREIGKEINMOGTNVDDPKIIYY APMRSLVQEMVGSFGKRLATYGITVAELTGDHQ LCKEEISATQIVCTPEKWDJITRKGGERTYTQLV RLIILDEHLLHDDRGFVLEALVARAIRNIEMTQE DVRLIGLSATLPNYEDVATFLRVDPAKGLFYFDN SRPPYPLEQTYVGTITEKAIKRRQIMNEITYCEKIM EHAGKNQVLVFVHSRKETGKTARARDMCLEKD TLGLFLREGSASTEVLRTEABCCKNLELKDLLPY GFAIHHAGMTRVDRTLVEDLFGDKHQVLVSTA TLAWGVNLPAHTVIIKGTQVYSPEKGRWTELGA LDLQMLGRAGRQYDTKGEGILITSHGELQYYL SLLNQQLPESQMYSKLPDMILNAEIVLGNVQNA KDAVNWLGYAYLYRMR.RSPTLYGISHDDLKGD PLLDQRRLDLVHTAALMLDKNNLVKYDKKTGN FQVTELGRIASHYYTNDTOYTYNQLKPTLSEIE LFRVFSLSSEFKNITVREEEKLELQKLLEERYPPVK ESIEEPSAKINVLLQAFISQLKLEGFALMADMYY VTQSAGRLMRAIFEIVLNRGWAQLTDKTLNLCK MIDKRMWQSMCPLRQFRKLPFEVVKKIEKKNIPP FERLYDLNIHNEIGELIRMPKMGKTIHKYVHLFPK LELSYHLQPITRSTLEVELTITPDFQWDEKVKHEKKNIPP FERLYDLNIHNEIGELIRMPKMGKTIHKYVHLFPK LELSYHLQPITRSTLEVELTITPDFQWDEKVHGSS EAFWILVEDVDSEVILHEYFILKAKY AQDEHLI TFFVYFFEPPPGYPTRVTOSPWILSCETQLEVSFR HLILFEKYPPTFELLDLQPLPYSALRNSAFESLYQ DKPFFPNFGTQVPTNTVYNSDENVLSCETQLEVSFR HLILFEKYPPTFELLDLQRLPYSALRNSAFESLYQ DKYPFPNFGTQVPTNTVYNSDENVLSCETQLEVSFR HLILFEKYPPTFELLDLGRLPSRWRYRSNQNINLFTVDVDEV HLIGGENGFVLEVICSRMYTSSQIERRRIVALSSS LSNAKGVAHVUGCSATSTTNFHNVRPPURDLEHI QGFNISHTQTKLLSMARPVFHAITKHSPKKPVTVF VPSRKQTRLTAADLQRGRFLHCTEKOL PYTLEKLSDSTLKETLLNOGYLHEGLSFMERRI VEQLFSSGAIQVVVASSLCWGMNVAAHLVIIM DTLYYNGKHHAVYDSPIYDVLQMVGHANRPLQ DDEGRCVIMCQGSKKDFFKKFLYEPLPVESHLD HCMHPHNAELTVIKTENKQDAVDYLTWFLLYR RMTQNPNYYNLQGISHRHLSDHLSELVEQTLSDL EQSKCISEDEMDVAYNLGUMGHAATYVINTTIEL FFMSLNAKTKVRCHLEISNAAAFYSNIPITHEEDN					RHNTLYFMSDAYMGCDQEYKFSVDVKEAETDS
	3619	A	3		DNIDETYGVNVQFESDEEGDEDVYGEVREAS DDDMEGDEAVVRCTLSANMYVDEILVWCASEL NIPEFFPLESPHKKVGYGLSSRTWLQGGGKVIEA GRDLLVASGELMSSKKKDLHPRDIDAFWLQRQL SRFYDDAIVSQKKADEVLEILKTASDDRECENQL VLLLGFNTFDFIKVLRQHRMMILYCTLLASAQSE AEKERIMGKMEADPELSKFLYQLHETEKEDLIRE ERSRERVRQSRMDTDLETMDLDQGGEALAPRQ VLDLEDLVFTQGSHFMANKRCQLPDGSFRRQRK GYEEVHVPALKPKPFGSEEQLLPVEKLPKYAQA GFEGFKTLNRIQSKLYRAALETDENLLLCAPTGA GKTNVALMCMLREIGKHINMDGTINVDDFKIIYI APMRSLVQEMVGSFGKRLATYGITVAELTGDHQ LCKEEISATQIIVCTPEKWDIITRKGGERTYTQLV RLIILDEIHLLHDDRGPVLEALVARAIRNIEMTQE DVRLIGLSATLPNYEDVATFLRVDPAKGLFYFDN SFRPVPLEQTYVGITEKKAIKRFQIMNEIVYEKIM EHAGKNQVLVFVHSRKETGKTARAIRDMCLEKD TLGLFLREGSASTEVLRTEAEQCKNLELKDLLPY GFAIHHAGMTRVDRTLVEDLFGDKHIQVLVSTA TLAWGVNLPAHTVIIKGTQVYSPEKGRWTELGA LDILQMLGRAGRPQYDTKGEGILITSHGELQYYL SLLNQQLPIESQMVSKLPDMLNAEIVLGNVQNA KDAVNWLGYAYLYIRMLRSPTLYGISHDDLKGD PLLDQRRLDLVHTAALMLDKNNLVKYDKKTGN FQVTELGRIASHYYITNDTVQTYNQLLKPTLSEIE LFRVFSLSSEFKNITVREEEKLELQKLLERVPIPVK ESIEEPSAKINVLLQAFISQLKLEGFALMADMVY VTQSAGRLMRAIFEIVLNRGWAQLTDKTLNLCK MIDKRMWQSMCPLRQFRKLPEEVVKKIEKKNFP FERLYDLNHNEIGELIRMPKMGKTIHKYVHLFPK LELSVHLQPITRSTLKVELTITPDFQWDEKVHGSS EAFWILVEDVDSEVILHHEYFLLKAKYAQDEHLI TFFVPVFEPLPPQYFIRVVSDRWLSCETQLPVSFR HLILPEKYPPPTELLDLQPLPVSALRNSAFESLYQ DKFPFFNPQTQTVFNTVYNSDDNVFVGAPTGSGK TICAEFAILRMLLQNSEGRCVYITPMRLWQEQVY MDWYEKFQDRLNKKVVLLTGETSTDLKLLGKG NIISTPEKWDILSRRWKQRKNVQNINLFVVDEV HLIGGENGPVLEVICSRMRYISSQIERPIRIVALSSS LSNAKDVAHWLGCSATSTFNFHPNVRPVPLELHI QGFNISHTQTRLLSMAKPVFHAITKHSPKKPVIVF VPSRKQTRLTAIDILTTCAADIQRQFFLHCTEKDL IPYLEKLSDSTLKETLLNGVGYLHEGLSPMERRL VEQLFSSGAIQVVVASRSLCWGMNVAAHLVIIM DTLYYNGKIHAYVDYPIYDVLQMVGHANRPLQ DDEGRCVIMCQGSKKDFFKKFLYEPLPVESHLD HCMHDHFNAEIVTKTIENKQDAVDYLTWFFLYR RMTQNPNYYNLQGISHRHLSDHLSELVEQTLSDL EQSKCISIEDEMDVAPLNLGMIAAYYYINYTTIEL FSMSLNAKTKVRGLIEIISNAAEYENIPIRHHEDN

C 1, C C C 1, C . C . C

SEQ ID	Method	Predicted	Predicted end	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid,
NO:		beginning	nucleotide	E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine,
		nucleotide location	location corresponding	I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine,
	ł	corresponding	to last amino	T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine,
		to first amino	acid residue of	X=Unknown, *=Stop codon, /=possible nucleotide deletion,
		acid residue of	peptide	\=possible nucleotide insertion
	ł	peptide sequence	sequence	
				SRMQLSAELQSDTEEILSKAIRLIQACVDVLSSNG
				WLSPALAAMELAQMVTQAMWSEDSYLRRLPPF
		1		PSGLFKRCTDKGVESVFDIMEMEDEERNALLQLT
•			İ	DSQIADVARFCNRYPNIELSYEVVDKDSIRSGGP
				VVVLVQLEREEEVTGPVIAPLFPQKREEGWWVV
		ĺ		IGDAKSNSLISIKRLTLQQKAKVKLDFVAPATGG
		}		RHNTLYFMSDAYMGCDQEYKFSVDVKEAETDS DSD
2620	ļ	1205	323	VIKMALAARLLPQFLHSRSLPCGAVRLRTPAVAE
3620	A	1203	323	VILIPSATLCYFCRCRLGLGAALFPRSARALAASA
			}	LPAQGSRWPVLSSPGLPAAFASFPACPQRSYSTE
				EKPQQHQKTKMIVLGFSNPINWVRTRIKAFLIWA
				YFDKEFSITEFSEGAKQAFAHVSKLLSQCKFDLL
			ł	EELVAKEVLHALKEKVTSLPDNHKNALAANIDEI
	ļ		ļ	VFTSTGDISIYYDEKGRKFVNILMCFWYLTSANIP
				SETLRGASVFQVKLGNQNVETKQLLSASYEFQR
				EFTQGVKPDWTIARIEHSKLLE
3621	A	2	2995	SSSRSRHSSISPVRLPLNSSLGAELSRKKKERAAA
				AAAAKMDGKESSYERSGSYSGRSPSPYGRRRSSS
				PFLSKRSLSRSPLPSRKSMKSRSRSPAYSRHSSSH
				SKKKRSSSRSRHSSISPVRLPLNSSLGAELSRKKK
	}	ļ		ERAAAAAAKMDGKESSYERSGSYSGRSPSPYG
				RRRSSSPFLSKRSLSRSPLPSRKSMKSRSRSPAYS RHSSSHSKKKRSSSRSRHSSISPVRLPLNSSLGAEL
				SRKKKERAAAAAAKMDGKESKGSPVFLPRKE
				NSSVEAKDSGLESKKLPRSVKLEKSAPDTELVNV
		İ		THLNTEVKNSSDTGKVKLDENSEKHLVKDLKAQ
				GTRDSKPIALKEEIVTPKETETSEKETPPPLPTIASP
)		PPPLPTTTPPPQTPPLPPLPPIPALPQQPPLPPSQPA
				FSQVPASSTSTLPPSTHSKTSAVSSQANSQPPVQV
				SVKTQVSVTAAIPHLKTSTLPPLPLPPLLPGDDDM
				DSPKETLPSKPVKKEKEQRTRHLLTDLPLPPELPG
				GDLSPPDSPEPKAITPPQQPYKKRPKICCPRYGER
				RQTESDWGKRCVDKFDIIGIIGEGTYGQVYKAKD
				KDTGELVALKKVRLDNEKEGFPITAIREIKILRQL
				IHRSVVNMKEIVTDKQDALDFKKDKGAFYLVFE
			•	YMDHDLMGLLESGLVHFSEDHIKSFMKQLMEGL
				EYCHKKNFLHRDIKCSNILLNNSGQIKLADFGLA
				RLYNSEESRPYTNKVITLWYRPPKLLLGEERYTP
				AIDVWSCGCILGELFTKKPIFQANLELAQLELISR LCGSPCPAVWPDVIKLPYFNTMKPKKQYRRRLR
	1			EEFSFIPSAALDLLDHMLTLDPSKRCTAEQTLQSD
	ļ			FLKDVELSKMAPPDLPHWQDCHELWSKKRRRQ
				RQSGVVVEEPPPSKTSRKETTSGTSTEPVKNSSPA
				PPQPAPGKVESGAGDAIGLADITQQLNQSELAVL
	1]		LNLLQSQTDLSIPQMAQLLNIHSNPEMQQQLEAL
	1			NOSISALTEATSQQQDSETMAPEESLKEAPSAPVI
	1			LPSAEQTTLEASSTPADMQNILAVLLSQLMKTQE
				PAGSLEENNSDKNSGPQGPRRTPTMPQEEAAGRS
				NGGNAL
3622	A	16	390	TPERGSAYPETAAVRRPAGECPITMSDLEAKLST
]			EHLGDKIKDEDIKLRVIGQDSSEIHFKVKMTTPLK
]		KLKKSYCQRQGVPVNSLRFLFEGQRIADNHTPEE
				LGMEEEDVIEVYQEQIGGHSTV
3623	A	2	1544	PPPAPGPDGLNEGCLHRLSMPHQRPRTCAMNPE

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CEOTO MALES D 1	ated Dung! - 1 - 3	Amino said saguence (A-Alenine C-Custaine B-Assett- 4-12
SEQ ID Method begin nucle locati	ning nucleotide otide location	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine,
to first acid to peptic	t amino acid residue of peptide sequence	X=Unknown, *=Stop codon, /=possible nucleotide deletion, \text{\tincrem{\text{\tincr{\text{\texiclex{\text{\text{\texi}\text{\text{\text{\text{\texicl
seque	nce	LTMESLGTLHGARGGGSGGGGGGGGGGGG
		GHEQELLASPSPHHARRGPRGSLRGPPPPPTAHQ
		ELGTAAAAAAASRSAMVTSMASILDGGDYRPE
		LSIPLHHAMSMSCDSSPPGMGMSNTYTTLTPLQP
		LPPISTVSDKFHHPHPHHHHHHHHHHHHHHQRLSGN
		VSGSFTLMRDERGLPAMNNLYSPYKEMPGMSQS
		LSPLAATPLGNGLGGLHNAQQSLPNYGPPGHDK
		MLSPNFDAHHTAMLTRGEQHLSRGLGTPPAAM MSHLNGLHHPGHTQSHGPVLAPSRERPPSSSSGS
		QVATSGQLEEINTKEVAQRITAELKRYSIPQAIFA
		QRVLCRSQGTLSDLLRNPKPWSKLKSGRETFRR
	į	MWKWLQEPEFQRMSALRLAACKRKEQEPNKDR
		NNSQKKSRLVFTDLQRRTLFAIFKENKRPSKEMQ
		ITISQQLGLELTTVSNFFMNARRRSLEKWQDDLS
3624 A 27	2152	TGGSSSTSSTCTKA SARKAEAATSGTAARDGSVGRNLVPPPSASAPK
JUZT A	2152	AEVESNEKDNRPEEEEQVIHEDDERPSEKNEFSR
		RKRSKSEDMDNVQSKRRRYMEEEYEAEFQVKIT
		AKGDINQKLQKVIQWLLEEKLCALQCAVFDKTL
		AELKTRVEKIECNKRHKTVLTELQAKIARLTKRF
	Ì	EAAKEDLKKRHEHPPNPPVSPGKTVNDVNSNNN
		MSYRNAGTVRQMLESKRNVSESAPPSFQTPVNT VSSTNLVTPPAVVSSQPKLQTPVTSGSLTATSVLP
		APNTATVVATTQVPSGNPQPTISLQPLPVILHVPV
		AVSSQPQLLQSHPGTLVTNQPSGNVEFISVQSPPT
		VSGLTKNPVSLPSLPNPTKPNNVPSVPSPSIQRNP
		TASAAPLGTTLAVQAVPTAHSIVQATRTSLPTVG
·		PSGLYSPSTNRGPIQMKIPISAFSTSSAAEQNSNTT PRIENQTNKTIDASVSKKAADSTSQCGKATGSDS
		SGVIDLTMDDEESGASQDPKKLNHTPVSTMSSSQ
		PVSRPLQPIQPAPPLQPSGVPTSGPSQTTIHLLPTA
	}	PTTVNVTHRPVTQVTTRLPVPRAPANHQVVYTT
		LPAPPAQAPLRGTVMQAPAVRQVNPQNSVTVRV
		PQTTTYVVNNGLTLGSTGPQLTVHHRPPQVHTEP
		PRPVHPAPLPEAPQPQRLPPEAGSTSRPSEATLEV SHAFRVKMAIVLVMECPGGGSKLCHC
3625 A 210	1115	ASPFLRPQGHDSGEREPFSQTPGLMQPFSIPVQIT
		LQGSRRRQGRTAFPASGKKRETDYSDGDPLDVH
		KRLPSSTGEDRAVMLGFAMMGFSVLMFFLLGTT
	•	ILKPFMLSIQREESTCTAIHTDIMDDWLDCAFTCG
		VHCHGQGKYPCLQVFVNLSHPGQKALLHYNEE
•	J	AVQINPKCFYTPKCHQDRNDLLNSALDIKEFFDH KNGTPFSCFYSPASQSEDVILIKKYDQMAIFHCLF
	[WPSLTLLGGALIVGMVRLTQHLSLLCEKYSTVV
		RDEVGGKVPYIEQHQFKLCIMRRSKGRAEKS
3626 A 9	921	SSVVEFSALSVSMACLSPSQLQKFQQDGFLVLEG
		FLSAEECVAMQQRIGEIVAEMDVPLHCRTEFSTQ
		EEEQLRAQGSTDYFLSSGDKIRFFFEKGVFDEKG
	1	NFLVPPEKSINKIGHALHAHDPVFKSITHSFKVQT LARSLGLQMPVVVQSMYIFKQPHFGGEVSPHQD
	l	ASFLYTEPLGRVLGVWIAVEDATLENGCLWFIPG
]	SHTSGVSRRMVRAPVGSAPGTSFLGSEPARDNSL
		FVPTPVQRGALVLIHGEVVHKSKQNLSDRSRQA
1 1		YTFHLMEASGTTWSPENWLQPTAELPFPQLYT
3627 A 231	644	INSSPRTGRDHQELNLHTERDSRSQRAVLKIPRQ

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SEQ ID NO:	Method	Predicted beginning nucleotide location corresponding to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methlonine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion
				NPGIFYWIFLPSRSHSASHGSRQRQVSCQGTQDEI LKMRNTFAELKNSLEALSSRMDQAEERIGTQAG VQWRDHGSLQPQPPEFKQCFHLSLPSSWDYRAC LS
3628	A	2	810	GCKHLLQNSWYDPRVREADRVGQRARRPRAAM DWLMGKSKAKPNGKKPAAEERKAYLEPEHTKA RITDFQFKELVVLPREIDLNEWLASNTTTFFHHIN LQYSTISEFCTGETCQTMAVCNTQYYWYDERGK KVKCTAPQYVDFVMSSVQKLVTDEDVFPTKYG REFPSSFESLVRKICRHLFHVLAHIYWAHFKETLA LELHGHLNTLYVHFILFAREFNLLDPKETAIMDD LTEVLCSGGRRGSTVGAVGMGPAAGAPGAQNH VKER
3629	A	699	1604	CSHGSSAVSAWSPLFQASEVERQLSMQVHALRE DFREKNSSTNQHIIRLESLQAEIKMLSDRKRELEH RLSATLEENDLLQGTVEELQDRVLILERQGHDKD LQLHQSQLELQEVRLSCRQLQVKVEELTEERSLQ SSAATSTSLLSEIEQSMEAEELEQEREQLTLLSVE MTALKEERDRLRVTSEDKEPKEQLQKAIRDRDE AIAKKNAVELELAKCRMDMMSLNSQLLDAIQQ KLNLSQQLEAWQDDMHRVIDRQLMDTHLKERS QPAAALCRGHSAGRGDEPSIAEGKRLFSFFRKI
3630	A	423	1	PAKVLTLDIYLSKTEGAQVDEPVVITPRAEDCGD WDDMEKRSSGRRSGRRRGSQKSTDSPGADAELP ESAARDDAVFDDEVAPNAASDNASAEKKVKSPR AALDGGVASAASPESKPSPGTKGQLRGESDRSK QPPPASSP
3631	A	2082	674	WSGFWQLPGVRGVGSAPGGDGAEFTSRRGSSRR PGAACPGCRGAGSERAPGGMGRRRAPELYRAPF PLYALQVDPSTGLLIAAGGGGAAKTGIKNGVHF LQLELINGRLSASLLHSHDTETRATMNLALAGDI LAAGQDAHCQLLRFQAHQQQGNKAEKAGSKEQ GPRQRKGAAPAEKKCGAETQHEGLELRVENLQA VQTDFSSDPLQKVVCFNHDNTLLATGGTDGYVR VWKVPSLEKVLEFKAHEGEIEDLALGPDGKLVT VGRDLKASVWQKDQLVTQLHWQENGPTFSSTP YRYQACRFGQVPDQPAGLRLFTVQIPHKRLRQPP PCYLTAWDGSNFLPLRTKSCGHEVVSCLDVSES GTFLGLGTVTGSVAIYIAFSLQCLYYVREAHGIV VTDVAFLPEKGRGPELLGSHETALFSVAVDSRCQ LHLLPSRRSVPVWLLLLLCVGLIIVTILLLQSAFPG FL
3632	A	942	40	PWCQRVEVRSCGSSKRSCSRWSGSSWDGSRSLG RGLNHTSLNRSPPFTPDTMTHCCSPCCQPTCCRT TCCRTTCWKPTTVTTCSSTPCCQPSCCVPSCCQP CCHPTCCQNTCCRTTCCQPTCVASCCQPSCCSTP CCQPTCCGSSCCGQTSCGSSCCQPICGSSCCQPCC HPTCYQTICFRTTCCQPTCCQPTCCRNTSCQPTCC GSSCCQPCCHPTCCQTICRSTCCQPSCVTRCCSTP CCQPTCGGSSCCSQTCNESSYCLPCCRPTCCQTT CYRTTCCRPSCCCSPCCVSSCCQPSCC
3633	A	605	3004	GPEGYRGRARHPSLGSTTGHCGGGRGAEGTGT DPAAPAARLNVDGLLVYFPYDYIYPEQFSYMRE LKRTLDAKGHGVLEMPSGTGKTVSLLALIMAYQ RAYPLEVTKLIYCSRTVPEIEKVIEELRKLLNFYE

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SEQ ID NO:	Method	Predicted beginning nucleotide location corresponding to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion
•				KQEGEKLPFLGLALSSRKNLCIHPEVTPLRFGKD VDGKCHSLTASYVRAQYQHDTSLPHCRFYEEFD AHGREVPLPAGIYNLDDLKALGRRQGWCPYFLA RYSILHANVVVYSYHYLLDPKIADLVSKELARK AVVVFDEAHNIDNVCIDSMSVNLTRRTLDRCQG NLETLQKTVLRIKETDEQRLRDEYRRLVEGLREA SAARETDAHLANPVLPDEVLQEAVPGSIRTAEHF LGFLRRLLEYVKWRLRVQHVVQESPPAFLSGLA QRVCIQRKPLRFCAERLRSLLHTLEITDLADFSPL TLLANFATLVSTYAKGFTIIIEPFDDRTPTIANPIL HFSCMDASLAIKPVFERFQSVIITSGTLSPLDIYPK ILDFHPVTMATFTMTLARVCLCPMIIGRGNDQVA ISSKFETREDIAVIRNYGNLLLEMSAVVPDGIVAF FTSYQYMESTVASWYEQGILENIQRNKLLFIETQ DGAETSVALEKYQEACENGRGAILLSVARGKVS EGIDFVHHYGRAVIMFGVPYVYTQSRILKARLEY LRDQFQIRENDFLTFDAMRHAAQCVGRAIRGKT DYGLMVFADKRFARGDKRGKLPRWIQEHLTDA NLNLTVDEGVQVAKYFLRQMAQPFHREDQLGL
3634	A	159	384	SLLSLEQLESEETLKRIEQIAQQL LKMSSKTASTNNIAQARRTVQQLRLEASIERIKV SKASADLMSYCEEHARSDPLLIGIPTSENPFKDKK TCIIL
3635	A	5	409	TELSQLEKAHPPADMGRRKSKRKPPPKKKMTGT LETQFTCPFCNHEKSCDVKMDRARNTGVISCTV CLEEFQTPITCILGNLGFFQRVGRGLESGPCSSGP LCALVQGQSRPEEQVPPSDFCGVRRCRAGFQCQ
3636	A	48	282	DHLKSCYQDSHEDPTKMKRFLFLLLTISLLVMVQ IQTGLSGQNDTSQTSSPSASSSMSGGIFLFFVANAI IHLFCFS
3637	A	1	1248	ARAGSVVGSAAARGPPAGCRCERAARLPSSPAR RRRCDWVEDGAGRMEILMTVSKFASICTMGAN ASALEKEIGPEQFPVNEHYFGLVNFGNTCYCNSV LQALYFCRPFREKGLAYKSQPRKKESLLTCLADL FHSIATQKKKVGVIPPKKFITRLRKENELFDNYM QQDAHEFLNYLLNTIADILQEERKQEKQNGRLPN GNIDNENNNSTPDPTWVHEIFQGTLTNETRCLTC ETISSKDEDFLDLSVDVEQNTSITHCLRGFSNTET LCSEYKYYCEECRSKQEAHKRMKVKKLPMILAL HLKRFKYMDQLHRYTKLSYRVVFPLELRLFNTS GDATNPDRMYDLVAVVVHCGSGPNRGHYIAIV KSHDFWLLFDDDIVEKIDAQAIEEFYGLTSDISKN SESGYILFYQSRD
3638	A	11	630	PAGIPVSTISSDRRASTDLTRKMKPDETPMFDPNL LKEVDWSQNTATFSPAISPTHPGEGLVLRPLCTA DLNRGFFKVLGQLTETGVVSPEQFMKSFEHMKK SGDYYVTVVEDVTLGQIVATATLIIEHKFIHSCAK RGRVEDVVVSDECRGKQLGNLLLSTLTLLSKKL NCYKITLECLPQNVGFYKKFGYTVSEENYMCRR FLK
3639	A	2	1200	PRVRLLRPSRSRSCRGLLSTRAPGPSPFRSLHSSPL LPHAMKSPFYRCQNTTSVEKGNSAVMGGVLFST GLLGNLLALGLLARSGLGWCSRRPLRPLPSVFY MLVCGLTVTDLLGKCLLSPVVLAAYAQNRSLRV LAPALDNSLCQAFAFFMSFFGLSSTLQLLAMALE

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				CWLSLGHPFFYRRHITLRLGALVAPVVSAFSLAF CALPFMGFGKFVQYCPGTWCFIQMVHEEGSLSV LGYSVLYSSLMALLVLATVLCNLGAMRNLYAM HRRLQRHPRSCTRDCAEPRADGREASPQPLEELD HLLLLALMTVLFTMCSLPVIYRAYYGAFKDVKE KNRTSEEAEDLRALRFLSVISIVDPWIFIIFRSPVFR IFFHKIFIRPLRYRSRCSNSTNMESSL
3640	A	930	182	PLPPPTLAMFLTRSEYDRGVNTFSPEGRLFQVEY AIEAIKLGSTAIGIQTSEGVCLAVEKRITSPLMEPS SIEKIVEIDAHIGCAMSGLIADAKTLIDKARVETQ NHWFTYNETMTVESVTQAVSNLALQFGEEDADP GAMSRPFGVALLFGGVDEKGPQLFHMDPSGTFV QCDARAIGSASEGAQSSLQEVYHKSMTLKEAIKS SLIILKQVMEEKLNATNIELATVQPGQNFHMFTK EELEEVIKDI
3641	A	2	1254	PTGQGGRRAEARSCLLSKAMLGRSGYRALPLGD FDRFQQSSFGFLGSQKGCLSPERGGVGTGADVPQ SWPSCLCHGLISFLGFLLLLVTFPISGWFALKIVPT YERMIVFRLGRIRTPQGPGMVLLLPFIDSFQRVDL RTRAFNVPPCKLASKDGAVLSVGADVQFRIWDP VLSVMTVKDLNTATRMTAQNAMTKALLKRPLR EIQMEKLKISDQLLLEINDVTRAWGLEVDRVELA VEAVLQPPQDSPAGPNLDSTLQQLALHFLGGSM NSMAGGAPSPGPADTVEMVSEVEPPAPQVGARS SPKQPLAEGLLTALQPFLSEALVSQVGACYQFNV VLPSGTQSAYFLDLTTGRGRVGHGVPDGIPDVV
3642		1	237	VEMAEADLRALLCRELRPLGAYMSGRLKVKGD LAMAMKLEAVLRALK RRGEIDMATEGDVELELETETSGPERPPEKPRKH
	A			DSGAADLERVTDYAEEKEIQSSNLETAMSVIGDR RSREQKAKQER
3643	A	94	541	RKERRRRRMEAVVFVFSLLDCCALIFLSVYFII TLSDLECDYINARSCCSKLNKWVIPELIGHTIVTV LLLMSLHWFIFLLNLPVATWNIYRYIMVPSGNM GVFDPTEIHNRGQLKSHMKEAMIKLGFHLLCFF MYLYSMILALIND
3644	A	95	2808	TSCRHFPITSEDPLNYLLILTVERIYAYQALPLGFL FCSRDPVPEYLNHCGVKYVLISDRASFCALHIFFS PFRNVFRPAAGGGIAPPPRLWFQPSLSDAEMEIPK LLPARGTLQGGGGGGIPAGGGRVHRGPDSPAGQ VPTRRLLPRGPQDGGPGRRREEASTASRGPGPS LFAPRPHQPSGGGGGGGDDFFLVLLDPVGGDVE TAGSGQAAGPVLREEAEEGPGLQGESGANPAG PTALGPRCLSAVPTPAPISAPGPAAAFAGTVTIHN QDLLLRFENGVLTLATPPPHAWEPGAAPAQQPG CLIAPQAGFPHAAHPGDCPELPPDLLLAEPAEPAP APAPEEEAEGPAAALGPRGPLGSGPGVVLYLCPE ALCGQTFAKKHQLKMHLLTHSSSQGQRPFKCPL GGCGWTFTTSYKLKRHLQSHDKLRPFGCPAEGC GKSFTTVYNLKAHMKGHEQENSFKCEVCEESFP TQAKLGAHQRSHFEPERPYQCAFSGCKKTFITVS ALFSHNRAHFREQELFSCSFPGCSKQYDKACRLK IHLRSHTGERPFLCDFDGCGWNFTSMSKLLRHKR KHDDDRRFMCPVEGCGKSFTRAEHLKGHSITHL STKPFVCPVAGCCARFSARSSLYIHSKKHLQDVD

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		corresponding to first amino acid residue of peptide sequence	to last amino acid residue of peptide sequence	T=Threonine, V=Vallne, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion
				TWKSRCPISSCNKLFTSKHSMKTHMVKRHKVGQ DLLAQLEAANSLTPSSELTSQRQNDLSDAEIVSLF
				SDVPDSTSAALLDTALVNSGILTIDVASVSSTLAG
]	}	HLPANNNNSVGQAVDPPSLMATSDPPQSLDTSLF FGTAATGFQQSSLNMDEVSSVSVGPLGSLDSLA
				MKNSSPEPQALTPSSKLTVDTDTLTPSSTLCENSV
İ				SELLTPAKAEWSVHPNSDFFGQEGETQFGFPNAA
3645	A	2194	1707	GNHGSQKERNLITVTGSSFLV TVSFHKTMASLKCSTVVCVICLEKPKYRCPACRV
3043	, a	2154	1707	PYCSVVCFRKHKEQCNPETRPVEKKIRSALPTKT
]				VKPVENKDDDDSIADFLNSDEEEDRVSLQNLKN
				LGESATLRSLLLNPHLRQLMVNLDQGEDKAKLM RAYMQEPLFVEFADCCLGIVEPSQNEES
3646	A	85	1948	ERGGGKAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAA
				APPWDDSGDDDEATTPADKSELHHTLKNLSLKL
İ				DDLSTCNDLIAKHGAALQRSLTELDGLKIPSESG
				EKLKVVNERATLFRITSNAMINACRDFLELAEIHS RKWQRALQYEQEQRVHLEETIEQLAKQHNSLER
}			j	AFHSAPGRPANPSKSFIEGSLLTPKGEDSEEDEDT
				EYFDAMEDSTSFITVITEAKEDSRKAEGSTGTSSA
		<u>.</u>		DWSSADNVLDGASLVPKGSSKVKRRVRIPNKPN YSLNLWSIMKNCIGRELSRIPMPVNFNEPLSMLQ
		}		RLTEDLEYHHLLDKAVHCTSSVEQMCLVAAFSV
				SSYSTTVHRIAKPFNPMLGETFELDRLDDMGLRS
		}		LCEQVSHHPPSAAHYVFSKHGWSLWQEITISSKF
				RGKYISIMPLGAIHLEFQASGNHYVWRKSTSTVH NIIVGKLWIDQSGDIEIVNHKTNDRCQLKFLPYSY
				FSKEAARKVTGVVSDSQGKAHYVLSGSWDEQM
				ECSKVMHSSPSSPSSDGKQKTVYQTLSAKLLWK
ļ]		KYPLPENAENMYYFSELALTLNEHEEGVAPTDS RLRPDQRLMEKGRWDEANTEKQRLEEKQRLSR
				RRRLEACGPGSSCSSEE
3647	A	46	5007	PTGDACVSTSCELASALSHLDASHLTENLPKAAS
				ELGQQPMTELDSSSDLISSPGKKGAAHPDPSKTS VDTGQVSRPENPSQPASPRVTKCKARSPVRLPHE
				GSPSPGEKAAAPPDYSKTRSASETSTPHNTRRVA
	ļ	,		ALRGAGPGAEGMTPAGAVLPGDPLTSQEQRQGA
)		PGNHSKALEMTGIHAPESSQEPSLLEGADSVSSR
				APQASLSMLPSTDNTKEACGHVSGHCCPGGSRE SPVTDIDSFIKELDASAARSPSSQTGDSGSQEGSA
				QGHPPAGAGGGSSCRAEPVPGGQTSSPRRAWAA
				GAPAYPQWASQPSVLDSINPDKHFTVNKNFLSN
				YSRNFSSFHEDSTSLSGLGDSTEPSLSSMYGDAE DSSSDPESLTEAPRASARDGWSPPRSRVSLHKED
]			PSESEEQIEICSTRGCPNPPSSPAHLPTQAAICPAS
				AKVLSLKYSTPRESVASPREKVACLPGSYTSGPD
			.	SSQPSSLLEMSSQEHETHADISTSQNHRPSCAEET TEVTSASSAMENSPLSKVARHFHSPPILSSPNMV
	1			NGLEHDLLDDETLNQYETSINAAASLSSFSVDVP
	1			KNGESVLENLHISESQDLDDLLQKPKMIARRPIM
				AWFKEINKHNQGTHLRSKTEKEQPLMPARSPDS
	1			KIQMVSSSQKKGVTVPHSPPQPKTNLENKDLSKK SPAEMLLTNGQKAKCGPKLKRLSLKGKAKVNSE
	1			APAANAVKAGGTDHRKPLISPQTSHKTLSKAVS
				QRLHVADHEDPDRNTTAAPRSPQCVLESKPPLAT

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				SGPLKPSVSDTSIRTFVSPLTSPKPVPEQGMWSRF HMAVLSEPDRGCPTTPKSPKCRAEGRAPRADSG PVSPAASRNGMSVAGNRQSEPRLASHVAADTAQ PRPTGEKGGNIMASDRLERTNQLKIVEISAEAVSE TVCGNKPAESDRRGGCLAQGNCQEKSEIRLYRQ VAESSTSHPSSLPSHASQAEQEMSRSFSMAKLAS SSSSLQTAIRKAEYSQGKSSLMSDSRGVPRNSIPG GPSGEDHLYFTPRPATRTYSMPAQFSSHFGREGH PPHSLGRSRDSQVPVTSSVVPEAKASRGGLPSLA NGQGIYSVKPLLDTSRNLPATDEGDIISVQETSCL VTDKIKVTRRHYCYEQNWPHESTSFFSVKQRIKS FENLANADRPVAKSGASPFLSVSSKPPIGRRSSGS IVSGSLGHPGDAAARLLRRSLSSCSENQSEAGTL LPQMAKSPSIMTLTISRQNPPETSSKGSDSELKKS LGPLGIPTPTMTLASPVKRNKSSVRHTQPSPVSRS KLQELRALSMPDLDKLCSEDYSAGPSAVLFKTEL EITPRRSPGPPAGGVSCPEKGGNRACPGGSGPKT SAAETPSSASDTGEAAQDLPFRRSWSVNLDQLLV SAGDQQRLQSVLSSVGSKSTILTLIQEAKAQSENE EÓVCFIVLNRKEGSGLGFSVAGGTDVEPKSITVH RVFSQGAASQEGTMNRGDFLLSVNGASLAGLAH GNVLKVLHQAQLHKDALVVIKKGMDQPRPSAR QEPPTANGKGLLSRKTIPLEPGIGRSVAVHDALC VEVLKTSAGLGLSLDGGKSSVTGDGPLVIKRVY KGGAAEQAGIIEAGDEILAINGKPLVGLMHFDA WNIMKSVPEGPVQLLIRKHRNSS
3648	A .	337	1564	KSRLSVTLMPVQLSEHPEWNESMHSLRISVGGLP VLASMTKAADPRFRPRWKVVLTFFVGAAILWLL CSHRPAPGRPPTHNAHNWRLGQAPANWYNDTY PLSPPQRTPAGIRYRIAVIADLDTESRAQEENTWF TYLKKGYLTFSDSGDKVAVEWDKDHGVLESHL AEKGRGMELSDLIVFNGKLYSVDDRTGVVYQIE GSKAVPWVILSDGDGTVEKGFKAEWLAVKDER LYVGGLGKEWTTTTGDVVNENPEWVKVVGYK GSVDHENWVSNYNALRAAAGIQPPGYLIHESAC WSDTLQRWFFLPRRASQERYSEKDDERKGANLL LSASPDFGDIAVSHVGAVVPTHGFSSFKFIPNTDD QIIVALKSEEDSGRVASYIMAFTLDGRFLLPETKI GSVKYEGIEFI
3649	A	1	775	PTRPGSGSAGGARVGSGEFGVEMAALAPLPPLPA QFKSIQHHLRTAQEHDKRDPVVAYYCRLYAMQ TGMKIDSKTPECRKFLSKLMDQLEALKKQLGDN EAITQEIVGCAHLENYALKMFLYADNEDRAGRF HKNMIKSFYTASLLIDVITVFGELTDENVKHRKY ARWKATYIHNCLKNGETPQAGPVGIEEDNDIEEN EDAGAASLPTQPTQPSSSSTYDPSNMPSGNYTGI QIPPGAHAPANTPAEVPHSTGVAK
3650	A	20	963	KMAATLGPLGSWQQWRRCLSARDGSRRLLLLL LLGSGQGPQQVGAGQTFEYLKREHSLSKPYQGE APRPCFLRDWELQVHFKIHGQGKKNLHGDGLAI WYTKDRMQPGPVFGNMDKFVGLGVFVDTYPNE EKQQERVFPYISAMVNNGSLSYDHERDGRPTEL GGCTAIVRNLHYDTFLVIRYVKRHLTIMMDIDGK HEWRDCIEVPGVRLPRGYYFGTSSITGDLSDNHD VISLKLFELTVERTPEEEKLHRDVFLPSVDNMKL

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				PEMTAPLPPLSGLALFLIVFFSLVFSVFAIVIGIILY NKWQEQSRKRFY
3651	A	1	1218	RSWAYVKKCKNNMCPNRGLHDGPEPCWLHHA AGTVSAVQARGLQPSQSRSRPRVPGLATALAYG PAHTPPLSRIGWAMQPPPPGPLGDCLRDWEDLQ QDFQNIQVSAAADAGSPPSRVSLAQGQGSGSPGC KPSLPAEAEGAAQELENQMKERQGLFFDMEAYL PKKNGLYLSLVLGNVNVTLLSKQAKFAYKDEYE KFKLYLTIILILISFTCRFLLNSRVTDAAFNFLLVW YYCTLTIRESILINNGSRIKGWWVFHHYVSTFLSG VMLTWPDGLMYQKFRNQFLSFSMYQSFVQFLQ YYYQSGCLYRLRALGERHTMDLTVEGFQSWMW RVLTFLLPFLFFGHFWQLFNALTLFNLAQDPQCK EWQVLMCGFPFLLLFLGNFFTTLRVVHHKFHSQ RHGSKKD
3652	A		164	VTTSCIPFAFGLGVRASERLAEIDMPYLLKYQPM MQTIGQKYCMDPAVIAGVLSRKSPGDKILVNMG DRTSMVQDPGSQAPTSWISESQVFQTTEVLTTRI TELQRRFPTWTPDQYLRGGLCAYSGGAGYVRSS QDLSCDFCNDVLARAKYLKRHGF
3653	A		909	IVRRDWQEVSDIHLAMANCKMTKSIRFPALEHC YTGGEVVLPKDQEEWKRRTGLLLYENYGQSETG LICATYWGMKIKPGFMGKATPPYDVQFHMEASV ENCIIVSMNTADPGSQGITHSLLLQVIDDKGSILPP NTEGNIGIRIKPVRPVSLFMCYEGDPEKTAKVEC GDFYNTGDRGKMDEEGYICFLGRSDDIINASGYR IGPAEVESALVEHPAVAESAVVGSPDPIRGEVVK AFIVLTPQFLSHDKDQLTKELQQHVKSVTAPYKY PRKVEFVSELPKTITGKIERKELRKKETGQM
3654	A	2	909	IVRRDWQEVSDIHLAMANCKMTKSIRFPALEHC YTGGEVVLPKDQEEWKRRTGLLLYENYGQSETG LICATYWGMKIKPGFMGKATPPYDVQFHMEASV ENCIIVSMNTADPGSQGITHSLLLQVIDDKGSILPP NTEGNIGIRIKPVRPVSLFMCYEGDPEKTAKVEC GDFYNTGDRGKMDEEGYICFLGRSDDIINASGYR IGPAEVESALVEHPAVAESAVVGSPDPIRGEVVK AFIVLTPQFLSHDKDQLTKELQQHVKSVTAPYKY PRKVEFVSELPKTITGKIERKELRKKETGQM
3655	A	2	2364	SPGPSLPESAESLDGSQEDKPRGSCAEPTFTDTG MVAHINNSRLKAKGVGQHDNAQNFGNQSFEEL RAACLRKGELFEDPLFPAEPSSLGFKDLGPNSKN VQNISWQRPKDIINNPLFIMDGISPTDICQGILGDC WLLAAIGSLTTCPKLLYRVVPRGQSFKKNYAGIF HFQIWQFGQWVNVVVDDRLPTKNDKLVFVHST ERSEFWSALLEKAYAKLSGSYEALSGGSTMEGL EDFTGGVAQSFQLQRPPQNLLRLLRKAVERSSL MGCSIEVTSDSELESMTDKMLVRGHAYSVTGLQ DVHYRGKMETLIRVRNPWGRIEWNGAWSDSAR EWEEVASDIQMQLLHKTEDGEFWMSYQDFLNN FTLLEICNLTPDTLSGDYKSYWHTTFYEGSWRTG SSAGGCRNHPGTFWTNPQFKISLPEGDDPEDDAE GNVVVCTCLVALMQKNWRHARQQGAQLQTIGF VLYAVPKEFQNIQDVHLKKEFFTKYQDHGFSEIF TNSREVSSQLRLPPGEYIIIPSTFEPHRDADFLLRV FTEKHSESWELDEVNYAEQLQEEKVSEDDMDQ

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				DFLHLFKIVAGEGKEIGVYELQRLLNRMAIKFKS FKTKGFGLDACRCMINLMDKDGSGKLGLLEFKI LWKKLKKWMDIFRECDQDHSGTLNSYEMRLVIE KAGIKLNNKVMQVLVARYADDDLIIDFDSFISCF LRLKTMFTFFLTMDPKNTGHICLSLEQVLGEGW EGICRIAPACPSTPPPPSSDVPGPASCPRLFPPWDL LPVSTVAADDHVGIEAL
3656	A	3	174	PLCTHYLLPELPEKSSRTSPRSRPGNMLSGDPHLP QPLCHCLDHCPCCFSGKRLVA
3657	A	1		DTRSTYHNAHSLPTYVKSPAPCQMTYIKSPAPCQ TQTCYVQGASPCQSYYVQAPASGSTSQYCVTDP CSAPCSTSYCCLAPRTFGVSPLRRWIQRPQNCNT GSSGCCENSGSSGCCGSGGCGCSCGCGSSGCCCL GIIPMKSRSPALL
3658	A	92	1537	SEAPVQPQPYTMTSFYSTSSCPLGCTMAPGARNV FVSPIDVGCQPVAEANAASMCLLANVAHANRVR VGSTPLGRPSLCLPPTSHTACPLPGTCHIPGNIGIC GAYGKNTLNGHEKETMKFLNDRLANYLEKVRQ LEQENAELETTLLERSKCHESTVCPDYQSYFRTIE ELQQKILCSKAENARLIVQIDNAKLAADDFRIKL ESERSLHQLVEADKCGTQKLLDDATLAKADLEA QQESLKEEQLSLKSNHEQEVKILRSQLGEKFRIEL DIEPTIDLNRVLGEMRAQYEAMVETNHQDVEQ WFQAQSEGISLQAMSCSEELQCCQSEILELRCTV NALEVERQAQHTLKDCLQNSLCEAEDRYGTELA
	-			QMQSLISNLEEQLSEIRADLERQNQEYQVLLDVK ARLENEIATYRNLTPLQSLFHACLLYFLSKLWPC HRWVSLWPWSQHGEMILKARVRRLRLVALGSG VPSPCPVFLQD
3659	A	2	402	DLLQCLNQLYSASTEMSCQQSQQQCQPPPKCTP KCPPKCPPKCPPKCPPQYSAPCPPPVSSCCG SSSGGCCSSEGGCCLSHHRPRQSLRRRPQSSSC CGSGSGQQSGGSSCCHSSGGSCCHSSGGCC
3660	A	26	710	CSAVEVKMAARTAFGAVCRRLWQGLGNFSVNT SKGNTAKNGGLLLSTNMKWVQFSNLHVDVPKD LTKPVVTISDEPDILYKRLSVLVKGHDKAVLDSY EYFAVLAAKELGISIKVHEPPRKIERFTLLQSVHI YKKHRVQYEMRTLYRCLELEHLTGSTADVYLEY IQRNLPEGVAMEVTKFCFFIFLDTIRTVTRTHQGA NLGNTIRRKRRKQVIKPQGGHFCLNLK
3661	A	2	370	DVSVAASEPTVYRNPTKMSCQQNQQQCQPPPKC PIPKYPPKCPSKCASSCPPPISSCCGSSSGGCCSSG GCGCCSSEGGGCCLSHHRHHRSHCHRPKSSNCY GSGSGQQSGGSGCCSGGGCC
3662	A	205	1277	RKSLPHPNPQKMLKKPLSAVTWLCIFIVAFVSHP AWLQKLSKHKTPAQPQLKAANCCEEVKELKAQ VANLSSLLSELNKKQERDWVSVVMQVMELESN SKRMESRLTDAESKYSEMNNQIDIMQLQAAQTV TQTSAGKETSPLRERGVPPHLQHCFYIPPDDFLGS PELEVFCDMETSGGGWTIIQRRKSGLVSFYRDW KQYKQGFGSIRGDFWLGNEHIHRLSRQPTRLRVE MEDWEGNLRYAEYSHFVLGNELNSYRLFLGNY TGNVGNDALQYHNNTAFSTKDKDNDNCLDKCA QLRKGGYWYNCCTDSNLNGVYYRLGEHNKHLD GITWYGWHGSTYSLKRVEMKIRPEDFKP

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3663	A	64	1456	LSSAKETLAQMYNTVWNMEDLDLEYAKTDINC GTDLMFYIEMDPPALPPKPPKPTTVANNGMNNN MSLQDAEWYWGDISREEVNEKLRDTADGTFLV RDASTKMHGDYTLTLRKGGNNKLIKIFHRDGKY GFSDPLTFSSVVELINHYRNESLAQYNPKLDVKL LYPVSKYQQDQVVKEDNIEAVGKKLHEYNTQFQ EKSREYDRLYEEYTRTSQEIQMKRTAIEAFNETIK IFEEQCQTQERYSKEYIEKFKREGNEKEIQRIMHN YDKLKSRISEIIDSRRRLEEDLKKQAAEYREIDKR MNSIKPDLIQLRKTRDQYLMWLTQKGVRQKKL NEWLGNENTEDQYSLVEDDEDLPHHDEKTWNV GSSNRNKAENLLRGKRDGTFLVRESSKQGCYAC SVVVDGEVKHCVINKTATGYGFAEPYNLYSSLK ELVLHYQHTSLVQHNDSLNVTLAYPVYAQQRR
3664	A	944	406	GATVEDQSCNFGSLRWVVSVPHISARSCPDPLLS RTGRVPGGRGAGLPRHHSPRCCLQVFFNGANVR QVDVPTLTGAFGILAAHVPTLQVLRPGLVVVHA EDGTTSKYFVSSGSIAVNADSSVQLLAEEAVTLD MLDLGAAKANLEKAQAELVGTADEATRAEIQIR IEANEALVKALE
3665	A	98	1388	ASQLAFGGKLTSTPSRDFQGCGRGAVTCCSFHEH RHQSGRCLSTGMAPNLKGRPRKKKPCPQRRDSF SGVKDSNNNSDGKAVAKVKCEARSALTKPKNN HNCKKVSNEEKPKVAIGEECRADEQAFLVALYK YMKERKTPIERIPYLGFKQINLWTMFQAAQKLG GYETITARRQWKHIYDELGGNPGSTSAATCTRR HYERLILPYERFIKGEEDKPLPPIKPRKQENSSQE NENKTKVSGTKRIKHEIPKSKKEKENAPKPQDAA EVSSEQEKEQETLISQKSIPEPLPAADMKKKIEGY QEFSAKPLASRVDPEKDNETDQGSNSEKVAEEA GEKGPTPPLPSAPLAPEKDSALVPGASKQPLTSPS ALVDSKQESKLCCFTESPESEPQEASFPRLPHHTG HRWQTRMRRRMTNCPPWQITLPTAP
3666	A	113	1492	LLQEMCTKTIPVLWGCFLLWNLYVSSSQTTYPGI KARITQRALDYGVQAGMKMIEQMLKEKKLPDL SGSESLEFLKVDYVNYNFSNIKISAFSFPNTSLAF VPGVGIKALTNHGTANISTDWGFESPLFVLYNSF AEPMEKPILKNLNEMLCPIIASEVKALNANLSTLE VLTKIDNYTLLDYSLISSPEITENYLDLNLKGVFY PLENLTDPPFSPVPFVLPERSNSMLYIGIAEYFFKS ASFAHFTAGVFNVTLSTEEISNHFVQNSQGLGNV LSRIAEIYILSQPFMVRIMATEPPIINLQPGNFTLDI PASIMMLTQPKNSTVETIVSMDFVASTSVGLVIL GQRLVCSLSLNRFRLALPESNRSNIEVLRFENILSS ILHFGVLPLANAKLQQGFPLPNPHKFLFVNSDIEV LEGFLLISTDLKYETSSKQQPSFHVWEGLNLISRQ WRGKSAP
3667	A	1	181	FRGRLGSGRNGGGSMNAPPAFESFLLFEGEKITIN KDTKVPNACLFTINKEDHTLGNIIK
3668	Α	212	431	VAGEAVPFFPMMYSEPLKPSYLALVLWYFLLTG YCITKPEVIFKIEQGEEPWILEKGFPSQCHPAKYL WCLHD
3669	A	458	1056	FSGVCFAGIAGSMATLLHDAVMNPAEVVKQRLQ MYNSQHRSAISCIRTVWRTEGLGAFYRSYTTQLT MNIPFQSIHFITYEFLQEQVNPHRTYNPQSHIISGG

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				LAGALAAAATTPLDVCKTLLNTQENVALSLANIS GRLSGMANAFRTVYQLNGLAGYFKGIQARVIYQ MPSTAISWSVYEFFKYFLTKRQLENRAPY
3670	A	145	298	RNPCPLTFLPSTLMVLLLSLTFFSALTFHSICQLRN TGVEVDIVFQRVSFL
3671	A	3	462	ILKVAKKERTMSSLPVPYKLPVSLSVGSCVIIKGT PIHSFINDPQLQVDFYTDMDEDSDIAFRFRVHFG NHVVMNRREFGIWMLEETTDYVPFEDGKQFELC IYVHYNEYEIKVNGHTHLRALSHRIPPSFVEDGC KCPRRYLPWTSVCVCN
3672	A	1	1028	HYAKLGTRPRLKFMSSPSLSDLGKREPAAAADE RGTQQRRACANATWNSIHNGVIAVFQRKGLPDQ ELFSLNEGVRQLLKTELGSFFTEYLQNQLLTKGM VILRDKIRFYEGQKLLDSLAETWDFFFSDVLPML QAIFYPVQGKEPSVRQLALLHFRNAITLSVKLED ALARAHARVPPAIVQMLLVLQGVHESRGVTEDY LRLETLVQKVVSPYLGTYGLHSSEGPFTHSCILEK RLLRRSRSGDVLAKNPVVRSKSYNTPLLNPVQE HEAEGAAAGGTSIRRHSVSEMTSCPEPQGFSDPP GQGPTGTFRSSPAPHSGPCPSRLYPTTQPPEQGLD PTRS
3673	A	2	712	RPPRVWYPELRELSAAAPRWSHRTAPGIMVFYF TSSSVNSSAYTIYMGKDKYENEDLIKHGWPEDI WFHVDKLSSAHVYLRLHKGENIEDIPKEVLMDC AHLVKANSIQGCKMNNVNVVYTPWSNLKKTAD MDVGQIGFHRQKDVKIVTVEKKVNEILNRLEKT KVERFPDLAAEKECRDREERNEKKAQIQEMKKR EKEEMKKKREMDELRSYSSLMKVENMSSNQDG NDSDEFM
3674	A	2	712	RPPRVWYPELRELSAAAPRWSHRTAPGIMVFYF TSSSVNSSAYTIYMGKDKYENEDLIKHGWPEDI WFHVDKLSSAHVYLRLHKGENIEDIPKEVLMDC AHLVKANSIQGCKMNNVNVVYTPWSNLKKTAD MDVGQIGFHRQKDVKIVTVEKKVNEILNRLEKT KVERFPDLAAEKECRDREERNEKKAQIQEMKKR EKEEMKKKREMDELRSYSSLMKVENMSSNQDG NDSDEFM
3675	A	921	1321	VTLAKMRVHISSCLKVQEQMANCPKFVPVVPTS QPIPSNIPNRSTFACPYCGARNLDQQELVKHCVE SHRSDPNRVVCPICSAMPWGDPSYKSANFLQHL LHRHKFSYDTFVDYSIDEEAAFQAALALSLSEN
3676	A		1856	TLGRWLLGVYETVAPTLACLPRPRLRRRRRRR RRMISRYTRKAVPQSLELKGITKHALNHHPPPEK LEEISPTSDSHEKDTSSQSKSDITRESSFTSADTGN SLSAFPSYTGAGISTEGSSDFSWGYGELDQNATE KVQTMFTAIDELLYEQKLSVHTKSLQEECQQWT ASFPHLRILGRQIITPSEGYRLYPRSPSAVSASYET TLSQERDSTIFGIRGKKLHFSSSYAHKASSIAKSSS FCSMERDEEDSIIVSEGIIEEYLAFDHIDIEEGFHG KKSEAATEKQKLGYPPIAPFYCMKEDVLAYVFD SVWCKVVSCMEQLTRSHWEGFASDDESNVAVT RPDSESSCVLSELHPLVLPRVPQSKVLYITSNPMS LCQASRHQPNVNDLLVHGMPLQPRNLSLMDKLL DLDDKLLMRPGSSTILSTRNWPNRAVEFSTSSLS YTVQSTRRRNPPPRTLHPISTSHSCAETPRSVEEIL

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				RGARVPVAPDSLSSPSPTPLSRNNLLPPIGTAEVE HVSTVGPQRQMKPHGDSSRAQSAVVDEPNYQQ PQERLLLPDFFPRPNTTQSFLLDTQYRRSCAVEYP HQARPGRGSAGPQLHGSTKSQSGGRPVSRTRQG P
3677	A	246	757	MRLQGAIFVLLPHLGPILVWLFTRDHMSGWCEG PRMLSWCPFYKVLLLVQTAIYSVVGYASYLVWK DLGGGLGWPLALPLGLYAVQLTISWTVLVLFFT VHNPGLALLHLLLLYGLVVSTALIWHPINKLAAL LLLPYLAWLTVTSALTYHLWRDSLCPVHQPQPT EKSD
3678	A	20	1508	RGKAEFFLAMAGTNALLMLENFIDGKFLPCSSYI DSYDPSTGEVYCRVPNSGKDEIEAAVKAAREAFP SWSSRSPQERSRVLNQVADLLEQSLEEFAQAESK DQGKTLALARTMDIPRSVQNFRFFASSSLHHTSE CTQMDHLGCMHYTVRAPVGVAGLISPWNLPLY LLTWKIAPAMAAGNTVIAKPSELTSVTAWMLCK LLDKAGVPPGVVNIVFGTGPRVGEALVSHPEVPL ISFTGSQPTAERITQLSAPHCKKLSLELGGKNPAII FEDANLDECIPATVRSSFANQGEICLCTSRIFVQK SIYSEFLKRFVEATRKWKVGIPSDPLVSIGALISK AHLEKVRSYVKRALAEGAQIWCGEGVDKLSLPA RNQAGYFMLPTVITDIKDESCCMTEEIFGPVTCV VPFDSEEVIERANNVKYGLAATVWSSNVGRVH RVAKKLQSGLVWTNCWLIRELNLPFGGMKSSGI GREGAKDSYDFFTEIKTITVKH
3679	A	1862	502	MAGTKPYMEIQTTIREYYEHLYANKLENLEEMD KFLDTYTLPRLNQEEVESLNRPITGSEIEAIINSLP TKKIPGPDRFTAKFYQRYKEELSNLIHYLGLSHH LLALNFIIVSFGKKSAWSSAQVKVTDTDFDGVEV RVFEGPPKPEEPLKRSVVYIHGGGWALASAKIRY YDELCTAMAEELNAVIVSIEYRLVPKVYFPEQIH DVVRATKYFLKPEVLQKYMVDPGRICISGDSAG GNLAAALGQQFTQDASLKNKLKLQALIYPVLQA LDFNTPSYQQNVNTPILPRYVMVKYWVDYFKG NYDFVQAMIVNNHTSLDVEEAAAVRARLNWTS LLPASFTKNYKPVVQTTGNARIVQELPQLLDARS APLIADQAVLQLLPKTYILTCEHDVLRDDGIMYA KRLESAGVEVTLDHFEDGFHGCMIFTSWPTNFSV GIRTRNSYIKWLDQNL
3680	A	249	2146	RSWGAPWFWRMRLLRRRHMPLRLAMVGCAFV LFLFLLHRDVSSREEATEKPWLKSLVSRKDHVLD LMLEAMNNLRDSMPKLQIRAPEAQQTLFSINQSC LPGFYTPAELKPFWERPPQDPNAPGADGKAFQK SKWTPLETQEKEEGYKKHCFNAFASDRISLQRSL GPDTRPPECVDQKFRRCPPLATTSVIIVFHNEAWS TLLRTVYSVLHTTPAILLKEIILVDDASTEEHLKE KLEQYVKQLQVVRVVRQEERKGLITARLLGASV AQAEVLTFLDAHCECFHGWLEPLLARIAEDKTV VVSPDIVTIDLNTFEFAKPVQRGRVHSRGNFDWS LTFGWETLPPHEKQRRKDETYPIKSPTFAGGLFSI SKSYFEHIGTYDNQMEIWGGENVEMSFRVWQC GGQLEIIPCSVVGHVFRTKSPHTFPKGTSVIARNQ VRLAEVWMDSYKKIFYRRNLQAAKMAQEKSFG DISERLQLREQLHCHNFSWYLHNVYPEMFVPDL

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				TPTFYGAIKNLGTNQCLDVGENNRGGKPLIMYS CHGLGGNQYFEYTTQRDLRHNIAKQLCLHVSKG ALGLGSCHFTGKNSQVPKDEEWELAQDQLIRNS GSGTCLTSQDKKPAMAPCNPSDPHQLWLFV
3681	A	2982	1869	LKDTLKSQMTQEASDEAEDMKEAMNRMIDELN KQVSELSQLYKEAQAELEDYRKRKSLEDVTAEY IHKAEHEKLMQLTNVSRAKAEDALSEMKSQYSK VLNELTQLKQLVDAQKENSVSITEHLQVITTLRT AAKEMEEKISNLKEHLASKEVEVAKLEKQLLEE KAAMTDAMVPRSSYEKLQSSLESEVSVLASKLK ESVKEKEKVHSEVVQIRSEVSQVKREKENIQTLL KSKEQEVNELLQKFQQAQEELAEMKRYSESSK LEEDKDKKINEMSKEVTKLKEALNSLSQLSYSTS SSKRQSQQLEALQQQVKQLQNQLAECKKQHQE VISVYRMHLLYAVQGQMDEDVQKVLKQILTMC KNQSQKK
3682	A	447	1024	AQALTAGRQLALAAPFIAPISPISLPRLNPPSQSW NSTPFFKVKLPPQKEVITSDELMAHLGNCLLSIKP QEKSEGLQLNFQQNVDDAMTVLPKLATGLDVN VRFTGVSDFEYTPECSVFDLLGIPLYHGWLVDPQ QSPEAVRAVGKLSYNQL/VGEDHHLQTLQ*HQP RDRKPDCRAVPGDHRGPSDLPRTV
3683	A	2	942	LEIKQEEKFVGQCIKEELMHGECVKEEKDFLKKE IVDDTKVKEEPPINHPVGCKRKLAMSRCETCGTE EAKYRCPRCMRYSCSLPCVKKHKAELTCNGVRD KTAYISIQQFTEMNLLSDYRFLEDVARTADHISR DAFLKRPISNKYMYFMKNRARRQGINLKLLPNG FTKRKENSTFFDKKKQQFCWHVKLQFPQSQA\ST *KKRVPDDKTINEILKPYIDPEKSDPVIRQRLKAYI RSQTGVQILMKIEYMQQNLVRYYELDPYKSLLD NLRNKVIIEYPTLHVVLKGSNNDMKVLHQVKSE STKNVGNEN
3684	A		1533	SLQENVQEKRVRVCPGLGGLLPNGTPSITAAAAP QVLWRHVQPGCSHHLHACVIRAACRAGEGHAD RHAGPPET/PVTLPSSWPWSSPWERQCPMH\L*AP GHAFRPVPTEHRRGWAALGHHRAAAGPLREPAS GSQPAPASC*PECHHGCPEQTRQCQDLLREAVV APEQRG*PCAHLQT*ATATTLCPQVPAGRVWQP GHSCHLLPHRHDGSH*HHCAAHRRPVTRRQAAH GVPLPDACYSPHHTLPAAPPPATRPAGHTATHPE *GGDLTPVPDGPHDCPRDVQGIPGAGGGSQLAPC CPPFPAAPVSVQGTQGLGPKNVLH*QWEGIRWQ KEPE/PGPPPEVELKRGAKCRIGDHGLGAVLGQG EYAS*SPSIPW*ASSSACPPLHPTP/TVYTQSPAAA PGWTRPPSP/PPPGLYPGP/PASHAPGVRGGISHQL YSLP*LCRECCSCP/PPPPAHGGRCPSLLPPEALAK LLL
3685	A	101	438	AWVLQCKINTELQTEVVMLKSMVLWLGEQVQS LQLQQQLHCHFNHTHICVTNLEYN\KEYPWDLV KAHLQGASTSNITFDIGELQKK\ILDLNKQTQEFQ PSL*AWTEFQQGLE
3686	Α .	105	845	VSDVVKNQLVEVQCRQDGCDAVENVHQMFMF NWFTDCLWTLFLSNYQPSVESSSPGGSATSDDHE FDPSADMLVHDFDDERTLEEEEMMEGETNFSSEI EDLAREGDMPIHELLSLYGYGSTVRLPEEDEEEE

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	-		·	EEEEEGEDDEDADNDDNSGCSGENKEENIKDSS GQEDETQSSNDDPSQSVASQDAQEIIRPRRCKYF DTNSEVEEESEEDEDYIP/SIISFFQSSDGI*SSSSSE DWKKEIMVGS
3687	A	49	1225	PVLVTSLRMREADTLRPPQLMEVSADIISTVEFN HTGELLATGDKGGRVVIFQREPESKNAPHSQGE YDVYSTFQSHEPEFDYLKSLEIEEKINKIKWLPQQ NAAHSLLSTNDKTIKLWKITERDKRPEGYNLKDE EGKLKDLSTVTSLQVPVLKPMDLMVEVSPRRIFA NGHTYHINSISVNSDCETYMSADDLRINLWHLAI TDRSFTP\NIVDIKPANMEDLTEVITASEFHPHHC NLFVYSSSKGSLRLCDMRAAALCDKHSKLFEEPE DPSNRSFFSEIIS\SVSDVKFSHSDRYMLTR\DYLT VKVWDLNMEARPIETYQVHDYLRSKLCSLYEND CIFDKFECAWNGSDR/IIMTGAYNNFFRMFDRNT KRDVTLEASRGSSKPRAVL
3688	A	1	401	KKVPGRLSEMSFSLNFTLPANTTSSPVT\DCGPSL GLAAGIPLLVATALLVALLFTLIHRRRSSIEAMEE SDRPCEISEIDDNPKISENPRRSPTHEKNTMGAQE AHIYVKTVAGSEEPVHDRYRPTIEMERRR
3689	A	698	889	GRVLVHCAMGVSRSATLVLAFLMIYENMTLVEA IPDGAGPPQISALTQAFVRQLQVLDNRLGRE
3690	A	61	153	MGAHLVRRYLGDASVEPDPLQMPTFPPDYGF
3691	A	61	153	MGAHLVRRYLGDASVEPDPLQMPTFPPDYGF
3692	A		2831	PLVRRLLRQTLRRVGGARAVREAVMRAVLTWR DKAEHCINDIAFKPDGTQLILAAGSRLLVYDTSD GTLLQPLKGHKDTVYCVAYAKDGKRFASGSAD KSVIIWTSKLEGILKYTHNDAIQCVSYNPITHQLA SCSSSDFGLWSPEQKSVSKHKSSSKIICCSWTNDG QYLALGMFNGIISIRNKNGEEKVKIERPGGSLSPI WSICWNPSSRWESFWMNRENEDAEDVIVNRYIQ EIPSTLKSAVYSSQGSEAEEEEPEEEDDSPRDDNL EERNDILAVADWG\QKVSFYQLSGKQIGKDRAL NFDPCCISYFTKGEYILLGGSDKQVSLFTKDGVR LGTVGEQNSWVWTGQAKPDSNYVVGGCQDGTI SFYQLIFSTVHGLYKDRYAYRDSMTDVIVQHLIT EQKVRIKCKELVKKIAIYRNRLAIQLPEKILIYELY SEDLSDMHYRVKEKIIKKFECNLLVVCANHIILC QEKRLQCLSFSGVKEREWQMESLIRYIKVIGGPP GREGLLVGLKNGQILKIFVDNLFAIVLLKQATAV RCLDMSASRKKLAVVDENDTCLVYDIDTKELLF QEPNANSVAWNTQCEDMLCFSGGGYLNIKASTF PVHRQKLQGFVVGYNGSKIFCLHVFSISAVEVPQ SAPMYQYLDRKLFKEAYQIACLGVTDTDWRELA MEALEGLDFETAKKERKKRGETNNDLFLADVFS YQGKFHEAAKLYKRSGHENLALEMYTDLCMFE YAKDFLGSGDPKETKMLITKQADWARNIKEPKA AVEMYISAGEHVKAIEICGDHGWVDMLIDIARK LDKAEREPLLLCATYLKKLDSPGYAAETYLKMG DLKSLVQLHVETQRWDEAFALGEKHPEFKDDIY MPYAQWLAENDRFEEAQKAFHKAGRQREAVQV LEQLTNNAVAESRFNDAAYYYWMLSMQCLDIA QDPAQKD
3693	A	3	1099	SSFPTCMRTVFHSNTSVSSLLHRPGHVTPQLTIHG GWRHHRDHTAIDEWDFNPSKFLIYTCLLLFSVLL

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-				PLRLDGIIQWSYWAVFAPIWLWKLLVVAGASVG AGVWARNPRYRTEGEACVEFKAMLIAVGIHLLL LMFEVLVCDRVERGTHFWLLVFMPLFFVSPVSV AACVWGFRHDRSLELEILCSVNILQFIFIALKLDRI IHWPWLVVFVPLWILMSFLCLVVLYYIVWSLLFL RSLDVVAEQRRTHVTMAISWITTVVPLLTFEVLL VHRLDGHNTFSYVSIFVPLWLSLLTLMATTFRRK GGNHWWFAIRRDF/CQDQLPQPTGKPPPPPLTDH HGEKALPLQNKDRGSWPASRGSPRLL
3694	A	483	761	PRSLIDYKSYMDTKLLVARFLEQSSCTMTPDIHE LVENIKSVLKSDEEHMEEAITSASFLEQIMAHSX QHIRAHKLPXETAGLXTSELRXLTP
3695	A	483	761	PRSLIDYKSYMDTKLLVARFLEQSSCTMTPDIHE LVENIKSVLKSDEEHMEEAITSASFLEQIMAHSX QHIRAHKLPXETAGLXTSELRXLTP
3696	A	456	733	LSAALWEEPILSLWSETKELTNRGKMNYPQIGPH RPHVKGLRVRPGPGTLSNAPKSLCPGMSNSDRGI H\GGEGQGPGKRAGHLGRGGGMSFL
3697	A	877	1873	VWL*TLS*HTCALMTVCRSCLVKYLEENNTCPT CRIVIHQSHPLQYIGHDRTMQDIVYKLVPGLQEA EMRKQREFYHKLGMEVPGDIKGETCSAKQHLDS HRNGETKADDSSNKEAAE
3698	A	1	572	KQCGIPHEVVRDENSSVYAEVSRLLLATGHWKR LRRDNPRFNLMLGERNRLPFGRLGHEPGLVQLV NYYRGADKLCRKASLVKLIKTSPELAESCTWFPE SYVIYPTNLKTPVAPAQNGIQPPISNSRTDEREFFL ASYNRKKEDGEGNVWIAKSSAGAKVWVQW*M TDLEEEIDIPSPVGLGLESEWPL
3699	A	2008	2432	LHCKMGALETQTHPCSQNMLRSLQKCCCKVEE HHLQPVQVLQTLLHSATAGTGCRRPARPPPAPPT PTPWRSRQSGKQSERAS*LKGRGRYGLGALGGR GGRALGGSRWPPPLPGETLFSGCKHRRRRGSD AAPGEEAGT
3700	A	33	1318	GYQIGMALASGPARRALAGSGQLGLGGFGAPRR GAYEWGVRSTRKSEPPPLDRVYEIPGLEPITFAG KMHFVPWLARPIFPPWDRGYKDPRFYRSPPLHE HPLYKDQACYIFHHRCRLLEGVKQALWLTKTKL IEGLPEKVLSLVDDPRNHIENQDECVLNVISHARL WQTTEEIPKRETYCPVIVDNLIQLCKSQILKHPSL ARRICVQNSTFSATWNRESLLLQVRGSGGARLST KDPLPTIASREEIEATKNHVLETFYPISPIIDLHECN IYDVKNDTGFQEGYPYPYPHTLYLLDKANLRPH RLQPDQLRAKMILFAFGSALAQARLLYGNDAKV LEQPVVVQSVGTDGRVFHFLVFQLNTTDLDSNE GVKNLAWVDSDQLLYQHFWCLPVIKKRVVVEP VGPVGFKPETFRKFLALYLHGAA
3701	A	86	465	WTLCGPEAGMVGYDPKPDGRNNTKFQVAVAGS VSGLVTRALISPFDVIKIRFQLQHERLSRSDPSAK YHGILQASRQILQEEGPTAFWKGHVPAQILSIGY GAVQFLSFEMLTELVHRGSVYDARE
3702	A	166	814	GFWEKTNQSSHSMDPLGAPSQFVDVDTLPSWGD SCQDELNSSDTTAEIFQEDTVRSPFLYNKDVNGK VVLWKGDVALLNCTAIVNTSNESLTDKNPVSESI FMLAGPDLKEDLQKLKGCRTGEAQLTKGFNLAA RFIIHTVGPKYKSRYRTAAESSLYSCYRNVLQLA

SEQ ID NO:	Method	Predicted beginning nucleotide location corresponding to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \ \=possible nucleotide insertion
				KEQSMSSVGFCVINSAKRGYPLKDATHIALRTVR RFLEIHGETIEKVV
3703	A	128	1255	SLGPSPKSATIPCCGDTMAPEEDAGGEALGGSFW EAGNYRRTVQRVEDGHRLCGDLVSCFQERARIE KAYAQQLADWARKWRGTVEKGPQYGTLEKAW HAFFTAAERLSALHLEVREKLQGQDSERVRAWQ RGAFHRPVLGGFRESRAAEDGFRKAQKPWLKRL KEVEASKKSYHAARKDEKTAQTRESHAKADSA VSQEQLRKLQERVERCAKEAEKTKAQYEQTLAE LHRYTPRYMEDMEQAFETCQAAERQRLLFFKD MLLTLHQHLDLSSSEKFHELHRDLHQGIEAASDE EDLRWWRSTHGPGMAMNWPQFEEWSLDTQRTI SRKEKGGRSPDEVTLTSIVPTRDGTAPPPQSPGSP GTGQDEEWSDEESP
3704	A	1	271	ARGEDLALATGGGPDTVTHSNMPCPNSLVYDC WLNIKECSVGEHTFEDLGLCPGRNQREKKRSYK DFLREEEKIAAQVRNSSKKKLKDSE
3705	A	170	1318	LNWANLVIMWPREEKEKVQDYSLGGLSPDLRI DVSRKKKILKAYDEDEDEDLYPDIHPPPSLPLPG QFTCPQCRKSFTRRSFRPNLQLANMVQIIRQMCP TPYRGNRSNDQGMCFKHQEALKLFCEVDKEAIC VVCRESRSHKQHSVLPLEEVVQEYKAKLQGHVE PLRKHLEAVQKMKAKEERRVTELKSQMKSELA AVASEFGRLTRFLAEEQAGLERRLREMHEAQLG RAGAAASRLAEQAAQLSRLLAEAQERSQQGGLR LLQDIKETFNRCEEVQLQPPEVWSPDPCQPHSHD FLTDAIVRKMSRMFCQAARVDLTLDPDTAHPAL MLSPDRRGVRLAERRQEVADHPKRFSADCCVLG AQGFRSGRHYWEVCMGP
3706	A		. 1996	SRERQTTWMDHNFAPAPPEMQSHGAPGPGTSFS HSHVLGRPIRPSRLPGGGSPLTPVLRKTIHLDTFP QSHIPQTSSRLGLGARTRSVPPQETGIALGASLSP LPTSSLVPRKLSSISLTLHQNSQARSLDRPLSHWE ELPTPGKKAAPHEGGRVSSPGSPPVTLVPGGRVH SEGPGNPGLTKSNRMLATEKPLVSSYLALPFQSR LAQSAPVLAEPGSLGQGHLVSVTDHMPTRASPG KGKPRARGIPRPRGRLQRANTTVNLTAMDTRTD AARHLATMATNRPSLAINLATPNTSQLDTGTEFP ALDIKLGTARDLSSVGTVKSGKTVNLATAGTIKP GTAMNLTTVGTTKPGMVMDLIASEPDKLGKAM ATRSTAKPDMTTEGIAMDSATSDPVKPDTITATV GTSRLETAMALARVNRAKLGTAKNSLALDTSR MGTAVGSVVPVTPDPATGKTTLGSVNNLTISDV ATCLLMPSRSTDLALDNTNAAMDRATEPASLDL ATEYKGKCRNLVGDGLGCREGEVCELGDGSMK PMSINSNLLGYIGIDTIIEQMRKKTMKTGFDFNIM VVGTEGCGAAAGLVAGSTKDPISFPQ
3707	A	3	549	SSSISRDFLGQAACASGTMLRWLRDFVLPTAACQ DAEQPMRYETLFQALDRNGDGVVDIGELQEGLR NLGIPLGQDAEEKIFTTGDVNKDGKLDFEEFMKY LKDHEKKMKLAFKSLDKNNDGKIEASEIVQSLQ TLGLTISEQQAELILQSIDVDGTMTVDWNEWRD YFLFNPVTDIEEIIR
3708	A	1	1866	EFRGAGRANMLAPRGAAVLLLHLVLQRWLAAG AQATPQVFDLLPSSSQRLNPGALLPVLTDPALND

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SEQ ID NO:	Method	Predicted beginning nucleotide location corresponding to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion
				LYVISTFKLQTKSSATIFGLYSSTDNSKYFEFTVM GRLSKAILRYLKNDGKVHLVVFNNLQLADGRRH RILLRLSNLQRGAGSLELYLDCIQVDSVHNLPRA FAGPSQKPETIELRTFQRKPQDFLEELKLVVRGSL FQVASLQDCFLQQSEPLAATGTGDFNRQFLGQM TQLNQLLGEVKDLLRQEVNETSFLRNTITECQAC GPLKFQSPTPSTVVPPASPAPPTRPPRRCDSNPCF RGVQCTDSRDGFQCGPCPEGYTGNGITCIDVDEC KYHPCYPGEHCINLSPGFRCDACPVGFTGPMVQ GVGISFAKSNKQVCTDIDECRNGACVPNSICVNT LGSYRCGPCKPGYTGDQIRGCKAERNCRNPELN PCSVNAQCIEERQGDVTCVCGVGWAGDGYICGK DVDIDSYPDEELPCSARNCKKDNCKYVPNSGQE DADRDGIGDACDEDADGDGILNEQDNCVLIHNV DQRNSDKDIFGDACDNCLSVLNNDQKDTDGDG RGDACDDDMDGDGIKNILDNCPKFPNRDQRDK DGDGVGDACDSCPDVSNPNQ
3709	A	144	417	TQAMEGLLHYINPAHAISLLSALNEERLKGQLCD VLLIVGDQKFRAHKNVLAASSEYFQSLFTNKENE SQTVFQLDFCEPDAFDNVLNYIY
3710	A	245	688	FGMLKNKGHSSKKDNLAVNAVALQDHILHDLQ LRNLSVADHSKTQVQKKENKSLKRDTKAIIDTGL KKTTQCPKLEDSEKEYVLDPKPPPLTLAQKLGLI GPPPPPLSSDEWEKVKQRSLLQGDSVQPCPICKE EFELRPQVFSIRG
3711	A	3	773	SLEMSSDGEPLSRMDSEDSISSTIMDVDSTISSGRS TPAMMNGQGSTTSSSKNIAYNCCWDQCQACFNS SPDLADHIRSIHVDGQRGGVFVCLWKGCKVYNT PSTSQSWLQRHMLTHSGDKPFKCVVGGCNASFA SQGGLARHVPTHFSQQNSSKVSSQPKAKEESPSK AGMNKRRKLKNKRRRSLARPHDFFDAQTLDAIR HRAICFNLSAHIESLGKGHSVVFHSTVSILLFFQIK YKTLQKNISTIISKSLKI
3712	A	2	344	RATWHNAGKEREAVQLMAGAEKRVKASHSFLR GLFGGNTRIEEACEMYTRAANMFKMAKNWSAA GNAFCQAAKLHMQLQSKHDSATSFVDAGNAYK KADPQGKTARHVACYLCV
3713	Α .	20	974	GAAATACSSSSSSSGAPATWAAHGPGKDVASPS SVSLSPRRSRLLVLRCGLRRNPERPSSSPALRRLL LLLLLLLLLLGFLLSPGPERGVGGGRFGRRLAL LWAAALGHVVSGKVMSRRAPGSRLSSGGGGG TNYSRSWNDWQPRTDSASADPGNLKYSSSRDRG GSSSYGLQPSNSAVVSRQRHDDTRVHADIQNDE KGGYSVNGGSGENTYGRKSLGQELRVNNVTSPE FTSVQHGSRALATKDMRKSQERSMSYCDESRLS YLLRRITRENDRDRRLATVKQLKEFIQQPENKLV LVKQLDILAAVHDVLNER
3714	A .	237	458	IFALKSPSYLLPCCTPEGKMDHKQLCWSHPQKSG QSSRSCCICSNQHGLIWKYSLNMCLQCCHQYVK DIGFIKL
3715	Α	970	1524	LCTLSPGISGTAGSCLTTEPGTELGTSFAQNGFYH EAVVLFTQALKLNPQDHRLFGNRSFCHERLGQP AWALADAQVALTLRPGWPRGLFRLGKALMGLQ RFREAAAVFQETLRGGSQPDAARELRSCLLHLTL QGQRGGICAPPLSPGALQPLPHAELAPSGLPSLRC

SEQ ID NO:	Method	Predicted beginning nucleotide location corresponding to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion
				PRSTALRSPGLSPLLH
3716	A	85	308	QGLPSTMVKLGCSFSGKPGKDPGDQDGAAMDS VPLISPLDISQLQPPLPDQVVIKTQTEYQLSSPDQQ NYTKSR
3717	A	58	618	GAGCTSPGLWARKAAARCLPTYPSRAQPSNVGR
	A			RRRRPGLGALAAGVPAMAESVERLQQRVQELE RELAQERSLQVPRSGDGGGGRVRIEKMSSEVVD SNPYSRLMALKRMGIVSDYEKIRTFAVAIVGVGG VGSVTAEMLTRCGIGKLLLFDYDKVELANMNRL FFQPHQAGLSKVQAAGHTPEE
3718	A .	3	593	RGAGGRAGGRADGQPNMADQRQRSLSTSGESL YHVLGLDKNATSDDIKKSYRKLALKYHPDKNPD NPEAADKFKEINNAHAILTDATKRNIYDKYGSLG LYVAEQFGEENVNTYFVLSSWWAKALFVFCGLL TCCYCCCCLCCCFNCCCGKCKPKAPEGEETEFY VSPEDLEAQLQSDEREATDTPIVIQPASATEP
3719	A		2173	SGGVRMGSRADGPRTSGHVTGKMAVFPWHSRN RNYKAEFASCRLEAVPLEFGDYHPLKPITVTESK TKKVNRKGSTSSTSSSSSSSVVDPLSSVLDGTDPL SMFAATADPAALAAAMDSSRRKRDRDDNSVVG SDFEPWTNKRGEILARYTTTEKLSINLFMGSEKG KAGTATLAMSEKVRTRLEELDDFEEGSQKELLN LTQQDYVNRIEELNQSLKDAWASDQKVKAPKN VHPGKLVYERIFSMCVDSRSVLPDHFSPENANDT AKETCLNWFFKIASIRELIPRFYVEASILKCNKFLS KTGISECLPRLTCMIRGIGDPL\GSVYARAYL\SRV GMEVAPHLKETLNKNFFDFLLTFKQIHGDTVQN QLVVQGVELPSYLPLYPPAMDWIFQCISYHAPEA LLTEMMERCKKLGNNALLLNSVMSAFRAEFIAT RSMDFIGMIKECDESGFPKHLLFRSLGLNLALAD PPESDRLQILNEAWKVITKLKNPQDYINCAEVWV EYTCKHFTKREVNTVLADVIKHMTPDRAFEDSY PQLQLIIKKVIAHFHDFSVLFSVEKFLPFLDMFQK ESVRVEVCKCI\RTPLSSINKSPPRTRSS*MPFCMF ARPCMTL/CNALTLEDEKRMLSYLINGFIKMVSF GRDFEQQLSFYVESRSMFCNLEPVLVQLIHSVNR LAMETRKVMKGNHSRKTAAFVRSWGAYWFITIP SLAGIFTRLNLYLHSG
3720	A	24	296	ENLFRAGFAFSLLRSSFYISKTYCSWFSNLISGSL ADFNSKGTRDYSPRQMAVRE/KVFDVIIRCFKRH GAEVIDTPVFELKVRNGQEETTW
3721	A	2	310	PSCLTCVGHCSIGGSCTMIGIMMPECHCSLHMTG PRCEEHVFILQQPGHIASILIPLLVLLLLALVAGVV FWHKRRVQGAKGFQHQRMTNGAMNVEIGNPTY K
3722	A	75	722	MELVAGCYEQVLFGFAVHPEPEACGDHEQWTL VADFTHHAHTASLSAVAVNSRFVVTGSKDETIHI YDMKKKIEHGALVHHSGTITCLKFYGNRHLISGA EDGLICIWDAKKWECLKSIKAHKGQVTFLSIHPS GKLALSVGTDKTLRTWNLVEGRSAFIKNIKQNA HIVEWSPRGEQYVVIIQNKIDIYQLDTASISGTITN EKRISSVKFLSES
3723	A	110	316	MELSDNRRSGGLEGLAEKCPNLTYLNLSGNKIK DLSTVEALVSGTVLSLDLLFLVKFSEICLCLLISI
3724	A	3	406	VDRGTEAWQRDPAFSGLQRVGGVDVSFVKGDS

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				VRACASLGVLSFPELEVVYEESRMVSLTAPYVSG FLAFREVPFLLELVQQLREKEPGLMPQVLLVDGN GVLHHRGFGVACHLGVLTDLPCVGVAKKLLQV DG
3725	A	3	406	VDRGTEAWQRDPAFSGLQRVGGVDVSFVKGDS VRACASLGVLSFPELEVVYEESRMVSLTAPYVSG FLAFREVPFLLELVQQLREKEPGLMPQVLLVDGN GVLHHRGFGVACHLGVLTDLPCVGVAKKLLQV DG
3726	A .	.1	433	SSDDRSLFRRLKLNYAIFDEGHMLKNMGSIRYQ HLMTINANNRLLLTGTPVQNNLLELMSLLNFVM PHMFSSSTSEIRRMFSSKTKSADEQSIYEKERIAH AKQIIKPFILRRVKEEVLKQLPPKKDRIELCAMSE KQEQLYLG
3727	A	6	383	RIPRGKACXTVLGRSTGELEGFASSRLPPQPCGW GQSSDLLSRIDLDELMKKDEPPLDFPDTLEGFEY AFNEKGQLRHIKTGEPFVFNYREHLHRWNQKRY EALGEIITKYVYELLEKDCNSKKVS
3728	A		2452	EIAGAAAENMLGSLLCLPGSGSVLLDPCTGSTISE TTSEAWSVEVLPSDSEAPDLKQEERLQELESCSG LGSTSDDTDVREVSSRPSTPGLSVVSGISATSEDIP NKIEDLRSECSSDFGGKDSVTSPDMDEITHDFLYI LQPKQHFQHIEAEADMRIQLSSSAHQLTSPPSQSE SLLAMFDPLSSHEGASAVVRPKVHYARPSHPPPD PPILEGAVGGNEARLPNFGSPMF*LPAEMEAFKQ RHS/YTPERLVRSRSS\DIVSSVRRPMSDPSWNRR P\GNEERELPPAAAIGATSLVAAPHSSSSSPSKDSS RGETEERKDSDDEKSDRNRPWWRKRFVSAMPK APIPFRKKEKQEKDKDDLGPDRFSTLTDDPSPRLS AQAQVAEDILDKYRNAIKRTSPSDGAMANYEST EVMGDGESAHDSPRDEALQNISADDLPDSASQA AHPQDSAFSYRDAKKKLRLALCSADSVAFPVLT\ HSTRNGLPDHTDPEDNEIVCFLKVQIAEAINLQD KNLMAQLQETMRCVCRFDNRTCRKLLASIAEDY RKRAPYIAYLTRCRQGLQTTQAHLERLLQRVLR DKEVANRYFTTVCVRLLLESKEKKIREFIQDFQK LTAADDKTAQVEDFLQFLYGAMAQDVIWQNAS EEQLQDAQLAIERSVMNRIFKLAFYPNQDGDILR DQVLHEHIQRLSKVVTANHRALQIPEVYLREAP WPSAQSEIRTISAYKTPRDKVQCILRMCSTIMNLL SLANEDSVPGADDFVPVLVFVLIKANPPCLLSTV QYISSFYASCLSGEESYWWMQFTAAVEFIKTIDD RK
3729	A	3	2452	EIAGAAAENMLGSLLCLPGSGSVLLDPCTGSTISE TTSEAWSVEVLPSDSEAPDLKQEERLQELESCSG LGSTSDDTDVREVSSRPSTPGLSVVSGISATSEDIP NKIEDLRSECSSDFGGKDSVTSPDMDEITHDFLYI LQPKQHFQHIEAEADMRIQLSSSAHQLTSPPSQSE SLLAMFDPLSSHEGASAVVRPKVHYARPSHPPPD PPILEGAVGGNEARLPNFGSPMF*LPAEMEAFKQ RHS/YTPERLVRSRSS\DIVSSVRRPMSDPSWNRR P\GNEERELPPAAAIGATSLVAAPHSSSSSPSKDSS RGETEERKDSDDEKSDRNRPWWRKRFVSAMPK APIPFRKKEKQEKDKDDLGPDRFSTLTDDPSPRLS AQAQVAEDILDKYRNAIKRTSPSDGAMANYEST

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SEO ID	Method	Predicted	Predicted end	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid,
NO:	MICTROG	beginning nucleotide	nucleotide location	E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine,
		location	corresponding	N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine,
	}	corresponding	to last amino	T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine,
		to first amino acid residue of	acid residue of peptide	X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion
		peptide sequence	sequence	t-possible addieonae insertion
		Sequence		EVMGDGESAHDSPRDEALQNISADDLPDSASQA AHPQDSAFSYRDAKKKLRLALCSADSVAFPVLT\
				HSTRNGLPDHTDPEDNEIVCFLKVOIAEAINLOD
		÷		KNLMAQLQETMRCVCRFDNRTCRKLLASIAEDY
		1	i	RKRAPYIAYLTRCRQGLQTTQAHLERLLQRVLR.
				DKEVANRYFTTVCVRLLLESKEKKIREFIQDFQK
				LTAADDKTAQVEDFLQFLYGAMAQDVIWQNAS
		ľ	[EEQLQDAQLAIERSVMNRIFKLAFYPNQDGDILR
				DQVLHEHIQRLSKVVTANHRALQIPEVYLREAP
	}			WPSAQSEIRTISAYKTPRDKVQCILRMCSTIMNLL
				SLANEDSVPGADDFVPVLVFVLIKANPPCLLSTV
				QYISSFYASCLSGEESYWWMQFTAAVEFIKTIDD
		L =		RK .
3730	Α	3	2452	EIAGAAAENMLGSLLCLPGSGSVLLDPCTGSTISE
				TTSEAWSVEVLPSDSEAPDLKQEERLQELESCSG
]		LGSTSDDTDVREVSSRPSTPGLSVVSGISATSEDIP
]	NKIEDLRSECSSDFGGKDSVTSPDMDEITHDFLYI
,				LQPKQHFQHIEAEADMRIQLSSSAHQLTSPPSQSE
	ļ			SLLAMFDPLSSHEGASAVVRPKVHYARPSHPPPD
	}			PPILEGAVGGNEARLPNFGSPMF*LPAEMEAFKQ RHS/YTPERLVRSRSS\DIVSSVRRPMSDPSWNRR
	l	1		P\GNEERELPPAAAIGATSLVAAPHSSSSSPSKDSS
				RGETEERKDSDDEKSDRNRPWWRKRFVSAMPK
				APIPFRKKEKQEKDKDDLGPDRFSTLTDDPSPRLS
				AQAQVAEDILDKYRNAIKRTSPSDGAMANYEST
				EVMGDGESAHDSPRDEALQNISADDLPDSASQA
				AHPQDSAFSYRDAKKKLRLALCSADSVAFPVLT\
		ļ		HSTRNGLPDHTDPEDNEIVCFLKVQIAEAINLQD
				KNLMAQLQETMRCVCRFDNRTCRKLLASIAEDY
		· ·		RKRAPYIAYLTRCRQGLQTTQAHLERLLQRVLR
		1		DKEVANRYFTTVCVRLLLESKEKKIREFIQDFQK
				LTAADDKTAQVEDFLQFLYGAMAQDVIWQNAS EEQLQDAQLAIERSVMNRIFKLAFYPNQDGDILR
				DQVLHEHIQRLSKVVTANHRALQIPEVYLREAP
		(WPSAQSEIRTISAYKTPRDKVQCILRMCSTIMNLL
			•	SLANEDSVPGADDFVPVLVFVLIKANPPCLLSTV
		1		QYISSFYASCLSGEESYWWMQFTAAVEFIKTIDD
				RK
3731	A	1	1305	VNTAMHEAKLMEECDELVEIIQQRKQMIAVKIK
				ETKVMKLRKLAQQVANCRQCLERSTVLINQAEH
				ILKENDQARFLQSAKNIAERVAMATASSQVLIPDI
				NFNDAFENFALDFSREKKLLEGLDYLTAPNPPSIR
				EELCTASHDTITVHWISDDEFSISSYELQYTIFTGQ
į				ANFISLYNSVDSWMIVPNIKQNHYTVHGLQSGTR YIFIVKAINQAGSRNSEPTRLKTNSQPFKLDPKMT
		·		HKKLKISNDGLQMEKDESSLKKSHTPERFSGTGC
		}		YVYGVLHNSDNS*MFISLSFPLSHRYAIGIAYKSA
				PKNEWIGKNASSWVFSRCNSNFVVRHNNKEML
				VDVPPHLKRLGVLLDYDNY/NMLSFYDPANSL\H
	}			LHTFDVTF\ILPVCPTFTIWNKSLMILSGLPAPDFI
	•			DYPERQECNCRPQESPYVSGMKTCH
3732	A	127	2832	LGQRLSLVPRPSLKRRLGKRLSLGLRERMMSLW
				WS/GPKVRTQATTGARPKTETKSVPAARPKTEAQ
				AMSGARPKTEVQVMGGARPKTEAQGITGARPKT
		l		DARAVGGARSKTDAKAIPGARPKDEAQAWAQS

		r =	<u></u>	
SEQ ID	Method	Predicted	Predicted end	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine,
NO:		beginning nucleotide	nucleotide location	E=Glutamic Acid, F=Fnenyialanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine,
]	ĺ	location	corresponding	N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine,
İ	ŀ	corresponding	to last amino	T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine,
	1	to first amino	acid residue of	X=Unknown, *=Stop codon, /=possible nucleotide deletion,
ł	ł	acid residue of	peptide	\=possible nucleotide insertion
		peptide sequence	sequence	
<u> </u>				EFGTEAVSQAEGVSQTNAVAWPLATAESGSVTK
	Ì		-	SK\ACLWIEN*SMWM/PETFPGTQGQKGIQPWFG
			ĺ	PGEETNMGSWCYSRPRAREEASNESGFWSADET
			}	STASSFWTGEETSVRSWPREESNTRSRHRAKHQT
	1	ļ		NPRSRPRSKQEAYVDSWSGSEDEASNPFSFWVG
		Ì		ENTNNLFRPRVREEANIRSKLRTNREDCFESESED
			ļ	EFYKQSWVLPGEEAN\IDSGTETKKILILPWKLRA
[{			QKDVDSDRVKQEPRFEEEVIIGSWFWAEKEASLE
		}		GGASAICESEPGTEEGAIGGSAYWAEEKSSLGAV
	1			AREEAKPESEEEAIFGSWFWDRDEACFDLNPCPV
· ·]			YKVSDRFRDAAEELNASSRPQTWDEVTVEFKPG
	ļ			LFHGVGFRSTSPFGIPEEASEMLEAKPKNLELSPE
	ł		•	GEEQESLLQPDQPSPEFTFQYDPSYRSVREIREHL
			ļ	RARESAESESWSCSCIQCELKIGSEEFEEFLLLMD
	ŀ	ļ	}	KIRDPFIHEISKIAMGMRSASQFTRDFIRDSGVVS
[['	LIETLLNYPSSRVRTSFLENMIHMAPPYPNLNMIE
		1		TFICQVCEETLAHSVDSLEQLTGNKGCFRHLTMT
		Į.		IDYHT\LIAN*YGPGFPLLF*PQAQCGETKFHVLK
	l			MLLNLSENPAVAKKLFSAKALSIFVGLFNIEETN
1				DNIQIVIKMFQNISNIIKSGKMSLIDDDFSLEPLISA
	İ			FREFEELAKQLQAQIDNQNDPEATGTTAFVGKG
				NNPSANRERLSPSVFCPGAQEAESLPARRVRGEE
		} .	ł	QRLLLEEVGARTADGIPEGW
3733	A	2	3274	DVPLIRIEEDTGEIFTTGARIDREKLCAGIPRDEHC
0.00	1			FYEVEVAILPDEIFRLVKIRFLIEDINDNAPLFPAT
		1		VINISIPENSAINSKYTLPAAVDPDVGINGVQNYE
		-		LIKSONIFGLDVIETPGGDKMPQLIVQKELDREEK
			1	DTYVMKVKVEDGGFPQRSSTAILQVSVTDTNDN
	İ)		HPVFKETEIEVSIPENAPVGTSVTQLHATDADIGE
				NAKIHFSFSNLVSNIARRLFHLNATTGLITIKEPLD
,				REETPNHKLLVLASDGGLMPARAMVLVNVTDV
	1			NDNVPSIDIRYIVNPVNDTVVLSENIPLNTKIALIT
				VTDKDADHNGRVTCFTDHEIPFRLRPVFSNQFLL
l		l		ETAAYLDYESTKEYAIKLLA\ADAGKPPLNQSAM
	[LFIKVKDENDNAPVFTQSFVTVSIPENNSPGIQLT
				KVSAMDADSGPNAKINYLLGPDAPPEFSLDCRT
	ŀ		ł	GMLTVVKKLDREKEDKYLFTILAKDNGVPPLTS
				NVTVFVSIIDQNDNSPVFTHNEYNFYVPENLPRH
1		ĺ		GTVGLITVTDPDYGDNSAVTLSILDENDDFTIDSQ
		}		TGVIRPNISFDREKQESYTFYVKAEDGGRVSRSSS
				AKVTINVVDVNDNKPVFIVPPSNCSYELVLPSTN
ĺ	1	1	{	PGTVVFQVIAVDNDTGMNAEVRYSIVGGNTRDL
]]	}	FAIDQETGNITLMEKCDVTDLGLHRVLVKANDL
				GQPDSLFSVVIVNLFVNESVTNATLINELVPQKH
		1		LKHQ*PQILEIADVSSPTSDYVKILVAAVAGTITV
}]	ļ	Į	VVVIFITAVVRCRQAPHLKAAQKNMQNSEWATP
				NPENROMIMMKKKKKKKKKHSPKNLLLNVVTIEE
1		1	1	TKADDVDSDGNRVTLDLPIDLEEQTMGKYNWV
]	ļ		TTPTTFKPDSPDLARHYKSASPQPAFQIQPETPLN
]	1]		LKHHIIQELPLDNTFVACDSISNCSSSSSDPYSVSD
1	ł	l	{	CGYPVTTFEVPVSVHTRPPVDLEVGGAQSGQVAI
				LTSSLMELLLCLMVAAFLPLELRPLGQQNVMSW
l	ļ	{	1	EQEAKILLVGYWGDGEWCHFHFHHLIPGPVNPG
I	1	į.	I	
1	L		1	VEDVOVIII DODOEDTODOGEI ODDUDDETII OIO
				YERKQYHILDSDSEDTQPSGELCPIPVRPFTILSIQ LLQDDGEHCGTKQGFQPAVQLGLLPHKTLK

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NO:	Method	Predicted beginning nucleotide location corresponding to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion
3734	A :	1	840	GTRPGHLPAPSDGFCV/HL*SIPSWGSF*GESL/EM QLITSLGLQEFDIARNVLELIYAQTLVWIGIFFCPL LPFIQMIMLFIMFYSKNISLMMNFQPPSKAWRAS QMMTFFIFLLFFPSFTGVLCTLAITIWRLKPSADC GPFRGLPLFIHSIYSWIDTLSTRPGYLWVVWIYRN LIGSVHFFFILTLIVLIITYLYWQITEGRKIMIRLLH EQIINEGKDKMFLIEKLIKLQDMEKKANPSSLVLE RREVEQQGFLHLGEHDGSLDLRSRRSVQEGNPR A
3735	A		432	VEVCRYLWKMTVDASQNVQCCVIFSHFFFIFN NLSKIKLLHTDTLLKIESKKHKAYLRSAAIEEERE SEFALRPTFDLTVRRNHLIEDVLNQLSQFENEDL RKELWVSFSGEIGYDLGGS/VKKEIFYCLFAEMIQ PEYGMFMY
3736	A	1542	343	KGAPSFVRLYQYPNFAGPHAALANKSFFKADKV TMLWNKKATAVLVIASTDVDKTGASYYGEQTL HYIATNGESAVVQLPKNGPIYDVVWNSSSTEFCA VYGFMPAKATIFNLKCDPVFDFGTGPRNAAYYS PHGHILVLAGFGNLILQI*AD/IMKVWNVKNYKLI SKPVASDSTYFAWCPDGEHILTATCAPRLRVNN GYKIWHYTGSILHKYDVPSNAELWQVSWQPFLD GIFPAKTITYQAVPSEVPNEEPKVATAYRPPALRN KPITNSKLHEEEPPQNMKPQSGNDKPLSKTALKN QRKHEAKKAAKQEARSDKSPDLAPTPAPQSTPR NTVSQSISGDPEIDKKIKNLKKKLKAIEQLKEQAA TGKQLEKNQLEKIQKETALLQELEDLELGI
3737	A	3190		VAMGTPRAQHPPPPQLLFLILLSCPWIQGLPLKEE EILPEPGSETPTVASEALAELLHGALLRRGPEMG YLPGPPLGPEGGEEETTTTIITTTTVTTTVTSPVLC NNNISEGEGYVESPDLGSPVSRTLGLLDCTYSIHV YPGYGIEIQVQTLNLSQEEELLVLAGGGSPGLAP RLLANSSMLGEGQVLRSPTNRLLLHFQSPRVPRG GGFRIHYQAYLLSCGFPPRPAHGDVSVTDLHPGG TATFHCDSGYQLQGEETLICLNGTRPSWNGETPS CMASCGGTIHNATLGRIVSPEPGGAVGPNLTCR WVIEAAEGRRLHLHFERVSLDEDNDRLMVRSGG SPLSPVIYDSDMDDVPERGLISDAQSLYVELLSET PANPLLLSLRFEAFEEDRCFAPFLAHGNVTTTDPE YRPGALATFSCLPGYALEPPGPPNAIECVDPTEPH WNDTEPACKAMCGGELSEPAGVVLSPDWPQSY SPGQDCVWGVHVQEEKRILLQVEILNVREGDML TLFDGDGPSARVLAQLRGPQPRRRLLSSGPDLTL QFQAPPGPPNPGLGQGFVLHFKEVPRNDTCPELP PPEWGWRTASHGDLIRGTVLTYQCEPGYELLGS DILTCQWDLSWSAAPPACQKIMTCADPGEIANG HRTASDAGFPVGSHVQYRCLPGYSLEGAAMLTC YSRDTGTPKWSDRVPKCALKYEPCLNPGVPENG YQTLYKHHYQAGESLRFFCYEGFELIGEVTITCV PGHPSQWTSQPPLCKVTQTTDPSRQLEGGNLAL AILLPLGLVIVLGSGVYIYYTKLQGKSLFGFSGSH SYSPITVESDFSNPLYEAGDTREYEVSI
3738	A	3190	664	VAMGTPRAQHPPPPQLLFLILLSCPWIQGLPLKEE EILPEPGSETPTVASEALAELLHGALLRRGPEMG YLPGPPLGPEGGEEETTTTIITTTTVTTTVTSPVLC NNNISEGEGYVESPDLGSPVSRTLGLLDCTYSIHV

SEQ ID NO:	Method	Predicted beginning nucleotide location corresponding to first amino acid residue of peptide	Predicted end nucleotide location corresponding to last amino acid residue of peptide, sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion
		sequence		YPGYGIEIQVQTLNLSQEEELLVLAGGGSPGLAP RLLANSSMLGEGQVLRSPTNRLLLHFQSPRVPRG GGFRIHYQAYLLSCGFPPRPAHGDVSVTDLHPGG TATFHCDSGYQLQGEETLICLNGTRPSWNGETPS CMASCGGTIHNATLGRIVSPEPGGAVGPNLTCR WVIEAAEGRRLHLHFERVSLDEDNDRLMVRSGG SPLSPVIYDSDMDDVPERGLISDAQSLYVELLSET PANPLLLSLRFEAFEEDRCFAPFLAHGNVTTTDPE YRPGALATFSCLPGYALEPPGPPNAIECVDPTEPH WNDTEPACKAMCGGELSEPAGVVLSPDWPQSY SPGQDCVWGVHVQEEKRILLQVEILNVREGDML TLFDGDGPSARVLAQLRGPQPRRRLLSSGPDLTL QFQAPPGPPNPGLGQGFVLHFKEVPRNDTCPELP PPEWGWRTASHGDLIRGTVLTYQCEPGYELLGS DILTCQWDLSWSAAPPACQKIMTCADPGEIANG HRTASDAGFPVGSHVQYRCLPGYSLEGAAMLTC YSRDTGTPKWSDRVPKCALKYEPCLNPGVPENG YQTLYKHHYQAGESLRFFCYEGFELIGEVTITCV PGHPSQWTSQPPLCKVTQTTDPSRQLEGGNLAL AILLPLGLVIVLGSGVYIYYTKLQGKSLFGFSGSH SYSPITVESDFSNPLYEAGDTREYEVSI
3739	A	734	445	LLEPEPAEEYTEQSEVEST/EGMILI*CCLYFAAFQ TNVSNIYFALQYVNRQFMAETQFTSGEKEQVDE WTVETVEVRVLCIAKLLSLSSVSNFYLY
3740	A	2	1578	MAHYITFLCMVLVLLLQNSVLAEDGEVRSSCRT APTDLVFILDGSYSVGPENFEIVKKWLVNITKNF DIGPKFIQVGVVQYSDYPVLEIPLGSYDSGEHLTA AVESILYLGGNTKTGKAIQFALDYLFAKSSRFLT KIAVVLTDGKSQDDVKDAAQAARDSKITLFAIG VGSETEDAELRAIANKPSSTYVFYVEDYIAISKIR EVMKQKLCEESVCPTRIPVAARDERGFDILLGLD VNKKVKKRIQLSPKKIKGYEVTSKVDLSELTSNV FPEGLPPSYVFVSTQRFKVKKIWDLWRILTIDG/* PQIAVTLNGVDKILLFTTTSVINGSQVVTFANPQV KTLFDEGWHQIRLLVTEQDVTLYIDDQQIENKPL HPVLGILINGQTQIGKYSGKEETVQFDVQKLRIY CDPEQNNRETACEIPGFCLNGPSDVGSTPAPCICP PGKPGLQGPKGDPGLPGNPGYPGQPGQDGKPVS TESLVISGISGITGYQGIAGTPGVPGSPGIQGARGL. PGYKGEPGRDGDK
3741	A	5048	1236	MSAPAGSSHPAASARIPPKFGGSAVSGAAAPAGP GAGPAPHQQNGPAQNQMQVPSGYGLHHQNYIA PSGHYSQGPGKMTSLPLDTQCGDYYSALYTVPT QNVTPNTVNQQPGAQQLYSRGPPAPHIVGSTLGS FQGAASSASHLHTSASQPYSSFVNHYNSPAMYS ASSSVASQGFPSTCGHYAMSTVSNAAYPSVSYPS LPAGDTYGQMFTSQNAPTVRPVKDNSFSGQNTA ISHPSPLPPLPSQQHHQQQSLSGYSTLTWSSPGLP STQDNLIRNHTGSLAVANNNPTITVADSLSCPVM QNVQPPKSSPVVSTVLSGSSGSSSTRTPPTANHPV EPVTSVTQPSELLQQKGVQYGEYVNNQASSAPT PLSSTSDDEEEEEEDEEAGVDSSSTTSSASPMPNS YDALEGGSYPDMLSSSASSPAPDPAPEPDPASAP APASAPAPVVPQPSKMAKPLAMAIQHFSLVIRML QHHLFLEYSPSNPVYSGFQQYPQQYPGVNQLSSS

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SEQ ID NO:	Method	Predicted beginning nucleotide location corresponding to first amino acid residue of	Predicted end nucleotide location corresponding to last amino acid residue of peptide	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion
	Ì	peptide	sequence	
		sequence		IGGLSLQSSPQPESLRPVNLTQERNILPMTPVWAP VPNLNADLKKLNCSPDSFRCTLTNIPQTQALLNK AKLPLGLLLHPFRDLTQLPVITSNTIVRCRSCRTYI NP\FVSFIDQRR*KCNLCYRVNDVPEEFMYNPLT RSYGEPHKRPEVQNS\TVEFIASSDYMLRPPQPAV YLFVLDVSHNAVEAGYLTI/LWCQSLLE\NLDKLP G\DSRT\RIGFMTFD\STYSFLQFTQEGLSQPQMLI VSDIDDVFLPTPDSLLVNLYESKELIKDLLNALPN MFTNTRETHSALGPALQAAFKLMSPTGGRVSVF QTQLPSLGAGLLQSREDPNQRSSTKVVQHLGPAT
				DFYKKLALDCSGQQTAVDLFLLSSQYSDLASLA CMSKYSAGCIYYYPSFHYTHNPSQAEKLQKDLK RYLTRKIGFEAVMRIRCTKGLSMHTFHGNFFVRS
	} .			TDLLSLANINPDAGFAVQLSIEESLTDTSLVCFQT ALLYTSSKGERRIRVHTLCLPVVSSLSDVYAGVD
				VQAAICLLANMAVDRSVSSSLSDARDALVNAVV DSLSAYGSTVSNLQHSALMAPSSLKLFPLYVLAL
				LKQKAFRTGTSTRLDDRVYAMCQIKSQPLVHLM KMIHPNLYRIDRLTDEGAVHVNDRIVPQPPLQKL
}				SAEKLTREGAFLMDCGSVFYIWVGKGCDNNFIE DVLGYTNFASIPQKMTHLPELDTLSSERARSFIT
1	!			WLRDSRPLSPILHIVKDESPAKAEFFQHLIEDRTE AAFSYYEFLLHVQQQICK
3742	A	934	68	SMLASQGVLLHPYGVPMIVPAAPYLPGLIQGNQE
				AAAAPDTMAQPYASAQFAPPQNGIPAEYTAPHP HPAPEYTGQTTVPEHTLNLYPPAQTHSEQSPADT
				SAQTVSGTRNKQD*RSTDGWPSPKTQTS*KHGK QVSSPSGLHVSNIPFR\FRDPDLRQMF\GQFGKILD
				VEIIFNERGSKGFGFVTFENSADADRAREK\LHGT VV\EGRKI\EVN\NATARVMTNKKTVNPYTNGWK
	Ì			LNPVVGAVYSPEFYAGTVLLCQANQEGSSMYSA PSTDFRGAKLHTSRPLLSGS
3743	A	3	1456	QFQQAWMQNKVPIPAPNEVLNDRKEDIKLEEKK
				KTQAEIEQEMATLQYTNPQLLEQLKIERLAQKQV EQIQPPPSSGTPLLGPQPFPGQGPMSQIPQGF/PTA
				PSISADANEHGS\KGPPGPQGQFRPPGPQGQMGP
				QGPPLHQGGGGPQGFMGPQGPQGPPQGLPRPQD
ĺ	[[MHGPQGMQRHPGPHGPLGPQGPPGPQGSSGPQG
				HMGPQGPPGPQGHIGPQGPPGPQGHLGPQGPPGT QGMQGPPGPRGMQGPPHPHGIQGGPGSQGIQGP
				VSQGPLMGLNPKGMQGPPGPRENQGPAPQGMI
	1			MGHPPQEMRGPHPPGGLLGHGPQEMRGPQEIRG
				MQGPPPQGSMLGPPQELRGPPGSQSQQGPPQGSL GPPPQGGMQGPPGPQGQQNPARGPHPSQGPIPFQ
			į	QQKTPLLGDGPRAPFNQEGQSTGPPPLIPGLGQQ
				GAQGRIPPLNPGQGPGPNKVS/ERGAPPRHEGRA PPRGRDGFPGPMKTLV
3744	A	1571	652	PLTGRKCPGWTHSGSRRSPRIAEEVPGFPKRAEA
				SRQFSETADRLELLRRAVMAAARATTPADGEEP
				APEAEALAAARERSSRFLSGLELVKQGAEARVFR GRFQGRAAVIKHRFPKGYRHPALEARLGRRRTV
				QEARALLRCRRAGISAPVVFFVDYASNCLYMEEI
{				EGSVTVRD\IFSPLWRLKKTPQGLSNLAKTIGQVL
				ARMHDEDLIHGDLTTSNMLLKPPLEQLNIVLIDF GLSFISALPEDKGVDLYVLEKAFLSTHPNTETVFE

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SEQ ID Method Predicted Predicted end Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, beginning nucleotide NO: I=Isoleucine, K=Lysine, L=Leucine, M=Methionine. nucleotide location N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, corresponding location T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, corresponding to last amino X=Unknown, *=Stop codon, /=possible nucleotide deletion, acid residue of to first amino \=possible nucleotide insertion acid residue of peptide sequence peptide sequence AFLKSYSTSSKKARPVLKKLDEVRLRGKKRSMV 127 1433 GSHRFSLASPLDPEVGPYCDTPTMRTLFNLLWLA 3745 A LACSPVHTTLSKSDAKKAASKTLLEKSQFSDKPV **QDRGLVVTDLKAESVVLEHRSYCSAKARDRHFA** GDVLGYVTPWNSHGYDVTKVFGSKFTQISPVWL QLKRRGREMFEVTGLHDVDQGWMRAVRKHAK GL\P*CLGSCLRTGLTMISG/YVLDSEDEIEELSKT VVQVAKNQHFDGFVVEVWNQLLSQKRVGLIHM LTHLAEALHQARLLALLVIPPAITPGTDQLGMFT HKEFEQLAPVLDGFSLMTYDYSTAHQPGPNAPL SWVRACVQVLDPKSKWRSKILLGLNFYGMDYA TSKDAREPVVGARYIQTLKDHRPRMVWDSQVSE HFFEYKKSRSGRHVVFYPTLKSLQVRLELARELG VGVSIWELGQGLDYFYDLL*VGIAASAVDVFFSK **PWSE** 898 IDRAAECRTKPLPMAVSIRGNADSIVACLVLMVL 3746 A 1 YLIKKRLVACAAVFYGFAVHMKIYPETYILPITL HLLPDRDNDKSLROFRYTFOACL*ELLKRLCNRT ALMFVAVAGLTFFALSFGFYYEYGWEFLEHTYF YHLTRRDIRHNFSPYFYMLYLTAESKWSFSLGIA AFLPQLILLSAVSFAYYRDLVFCWFLHTSIFVTFN KVCTSQYFLWYLCLLPLVMPLVRMPWKRAVVL LMLWFIGQAMWLAPAYVLEFQGKNTFLFIWLA **GLFFLLINCSILIQUISHYKEEPLTERIKYD** 3747 A 2325 MVISFQGLVTFGDVAVDFSQEEWEWLNPIQRNL 1 YRKVMLENYRNLASLGLCVSKPDVISSLEQGKEP WTVKRKMTRAWCPDLKAVWKIKELPLKKDFCE GKLSQAVITERLTSYNLEYSLLGEHWDYDALFET **QPGLVTIKNLAVDFRQQLHPAQKNFCKNGIWEN** NSDLGSAGHCVAKPDLVSLLEQEKEPWMVKREL TGSLFSGQRSVHETQELFPKQDSYAEGVTDRTSN TKLDCSSFRENWDSDYVFGRKLAVGQETQFRQE PITHNKTLSKERERTYNKSGRWFYLDDSEEKVH NRDSIKNFQKSSVVIKQTGIYAGKKLFKCNECKK TFTOSSSLTVHQRIHTGEKPYKCNECGKAFSDGS SFARHQRCHTGKKPYECIECGKAFIQNTSLIRHW RYYHTGEKPFDCIDCGKAFSDHIGLNQHRRIHTG EKPYKCDVCHKSF\RYGSSLTVHQRIHTGEKPYE CDVCRKAFSHHASLT\Q\HQRVHSGEKPFKCKEC GKAFRONIHLASHLRIHTGEKPFECAECGKSFSIS SQLATHQRIHTGEKPYECKVCSKAFTQKAHLAQ HQKTHTGEKPYECKECGKAFSQTTHLIQHQRVH TGEKPYKCMECGKAFGDNSSCTQHQRLHTGQRP YECIECGKAFKTKSSLICHRRSHTGEKPYECSVC GKAFSHRQSLSVHQRIHSGKKPYECKECRKTFIQI GHLNQHKRVHTGERSYNYKKSRKVFRQTAHLA **HHQRIHTGESSTCPSLPSTSNPVDLFPKFLWNPSS** LPSP 3748 823 ī **GGYTKSGYDSACKDFVPHDLEVQIPGRVFLVTG** A GNSGIGKATALEIAKRGGTVHLVCRDQAPAEDA RGEIIRE\SGNQNIFLHIVDLSDPKKIWKFVENFKQ EHKLHVL\VNNAGCMVNKREAHKKMDFEKNFG COYSGVCTFLTTRPDPLCWRKNTDPRVIT\VSSG GMLVQKLNNQ*SPVRKNTIWMGTMVYAQNKVS

SEQ ID NO:	Method	Predicted beginning nucleotide location corresponding to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon,/=possible nucleotide deletion, \possible nucleotide insertion
				ERQQVVLT\ERWGPRAPG\IHFSSMHPGWA\DTPG VRQAMPGFHVQASGYRLRSEAQGADTMLWLAL SSARSRTAQRP
3749	A	1939	715	GFLRLSQAT\RQRLSIPVMVLTLDPTRD\QCFGDR FSRLLLDEFLGYDDIL\MSSVKGLAENEENKGFLR NVVSGEHYRFV\SMWMART\SYLAAFANHGQSF TLSVSHACCGYSHHQIFVFIVDLLQMLEMNMAIA FPAAPLLTVILALVGMEAIMSEFFNDTTTAFYIILI VWLADQYDAICCHTSTSKRHWLRFFYLYHFAFY AYHYRFNGQYSSLALVTSWLFIQHSMIYFFHHYE LPAILQHVRIQ\EMLLQAPTLGPGTPTA\LPDDMN NNSGAPATAP\DSAGQPPALGPVSPGASGSPGPV AAAPSSLVAAAASVAAAAGGDLGWMAETAAIIT DASFLSGLSASLLERRPASPLGPAGGLPHAPQDS VPPSDSAASDTTPLGAAVGGPSPASMAPTEAPSE VGS
3750	A	2	844	GLLEPFSKLLSFVIQNAVFTLAYLVELCGLCYRA FTKERDKFYLSRSVVLELLQALKLKSPLPDTNLL LLVQFICADAGTKLAESTILSKQMIASVPGCGTA AMECVRQYINEVLDFM\ADMHTLTKLKSHMKTC SQPLHEDTFGGHLKVGLAQIAAMDISRGNHRDN KAVIRYLPWLYHPPSAMQQGPKEFIECVSHIRLL SWLLLGSLTHNAVC/LKWPPLPGLPIPLDAGSHV ADHLIVILIGFPEQSKTSVL\HMCSLFHAF\SLAQL WDSLLARQSGRW
3751	A	431	2	AFTRKCEETAFIVPQCEIIPTE/WVCRRIPTGSSLER NPGVKEGCEFCPPKVEMFFKDDANHDPQWSRQ QLIAAKFGFAALGI/QTEVDIMSHAT*AVFEIPEKS RL\PQNCTPVDMKIEFGVHVTSKEILTDVIDNDS* RHSPS
3752	A	131	1278	AWSGSGLLVLCINTASMPMISVLGKMFLWQREG PGGRWTCQTSRRVSSDPAWAVEWIELPRGLSLSS LGSARTLRGWSRSSRPSSVDSQDLPEVNVGDTV AMLPKSRRALTIQEIAALARSSLHGISQVVKDHV TKPTAMAQGRVAHLIEWKGWSKPSDSPAALESA FSSYSDLSEGEQEARFAAGVAEQFAIAEAKLRA WSSVDGEDSTDDSYDEDFAGGMDTDMAGQLPL GPHLQDLFTGHRFSRPVRQGSVEPESDCSQTVSP DTLCSSLCSLEDGLLGSPARLA\PSCWAMSCFSPN CPPAGKVPSAAW/APLEAQDSLYNSPLTESCLSP AEEEPAPCKDCQPLCPPLTGSWERQRQASDLASS GVVSLDEDEAEPEQ
3753	A	2	3338	YYSSVRQRVTCEEPRFRECAAALIEGSATEVYAG EWRADRRSGFGVSQRSNGLRYEGEWLGNRRHG YGRTTRPDGSREEGKYKRNRLVHGGRVRSLLPL ALRRGKVKEKVDRAVEGARRAVSAARQRQEIA AARAADALLKAVAASSVAEKAVEAARMAKLIA QDLQPMLEAPGRRPRQDSEGSDTEPLDEDSPGV YENGLTPSEGSPELPSSPASSRQPWRPPACRSPLP PGGDQGPFSSPKAWPEEWGGAGAQAEELAGYE AEDEAGMQGPGPRDGSPLLGGCSDSSGSLREEE GEDEEPLPPLRAPAGTEPEPIAMLVLRGSSSRGPD AGCLTEELGEPAATERPAQPGAANPLVVGAVAL LDLSLAFLFSQLLT SSLLEKMTSSDKDFRFMATSDLMSELQKDSIQLD

SEQ ID Method Predicted Predicted end Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E-Glutamic Acid, F-Phenylalanine, G-Glycine, H-Histidine, nucleotide beginning NO: nucleotide location I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, corresponding location T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, corresponding to last amino to first amino acid residue of X=Unknown, *=Stop codon, /=possible nucleotide deletion, acid residue of >=possible nucleotide insertion peptide peptide sequence sequence EDSERKVVKMLLRLLEDKNGEVQNLAVKWLGV PLGAFHASLLHCLLPQLSSPRLAVRKRAVGALGH LATACSTDLFVELADHLLDRLPGPRVPTSPTAIRT LIQCLGSVGRQAGHRLGAHLDRLVPLVEDFCNL **DDDELRESCLQAFEAFLRKCPKEMGPHVPNVTS** LCLQYIKHDPNYNYDSDEDEEQMETEDSEFSEQE SEDEYSDDDDMSWKVRRAAAKCIAALISSRPDL LPDFHCTLAPVLIRRFKEREENVKADVFTAYIVL LRQTRPPKGWLEAMEEPTQTGSNLHMLRGQVPL VVKALQRQLKDRSVRARQGCFSLLTELAGVLPG SLAEHMPVLVSGIIFSLADRSSSSTIRMDALAFLQ GLLGTEPAEAFHPHLPILLPPVMACVADSFYKIA **AEALVVLQELVRALWPLHRPRMLDPEPYVGEMS** AVTLARLRATDLDQEVKERAISCMGHLVGHLGD RLGDDLEPTLLLLLDRLRNEITRLPAIKALTLVAV SPLQLDLQPILAEALHILAŞFLRKNQRALRLATLA ALDALAQSQGLSLPPSAVQAVLAELPALVNESD MHVAQLAVDFLATVTQAQPASLVEVSGPVLSEL LRLLRSPLLPAGVLAAAEGFLQALVGTRPPCVDY AKLISLLTAPVYEQAVDGGPGLHKQVFHSLARC VAALSAACPQ\EAESTASRLVCDARSPHSSTGVK VLAFLSLAEVGQVAGPGHERELKAVLLEALGSPS **EDVRAAASYALGRVGAGSLPDFLPFLLEQIEAEP** RRQYLLLHSLKEALGAAQPDSLKPYAEDIWALL FORCEGAEEGTRGVVAECIGKLVLVNPSFLLPRL RKQLAAGRPHTRSTVITAVKFLISDQPHPIDPLLK SFIAVHNKPSLVRDLLDDILPLLYQETKIRRDLIRE VEMGPFKHTVDDGLDVRKAAFECMYSLLESCLG **QLDICEFLNHVEDGLKDHYDIRMLTFIMVARLAT** LCPAPVLQRVDRLIEPLRATCTAKVKAGSVKQEF **EKQDELKRSAMRAVAALLTIPEVGKSPIMADFSS QIRSNPELAALFESIQKDSTSAPSTDSMELS** 3755 3338 SSLLEKMTSSDKDFRFMATSDLMSELQKDSIQLD Α 2 EDSERKVVKMLLRLLEDKNGEVQNLAVKWLGV PLGAFHASLLHCLLPQLSSPRLAVRKRAVGALGH LATACSTDLFVELADHLLDRLPGPRVPTSPTAIRT LIQCLGSVGRQAGHRLGAHLDRLVPLVEDFCNL DDDELRESCLQAFEAFLRKCPKEMGPHVPNVTS LCLQYIKHDPNYNYDSDEDEEQMETEDSEFSEQE SEDEYSDDDDMSWKVRRAAAKCIAALISSRPDL LPDFHCTLAPVLIRRFKEREENVKADVFTAYIVL LROTRPPKGWLEAMEEPTQTGSNLHMLRGQVPL VVKALQRQLKDRSVRARQGCFSLLTELAGVLPG SLAEHMPVLVSGIIFSLADRSSSSTIRMDALAFLQ GLLGTEPAEAFHPHLPILLPPVMACVADSFYKIA **AEALVVLQELVRALWPLHRPRMLDPEPYVGEMS** AVTLARLRATDLDQEVKERAISCMGHLVGHLGD RLGDDLEPTLLLLLDRLRNEITRLPAIKALTLVAV SPLQLDLQPILAEALHILASFLRKNQRALRLATLA ALDALAQSQGLSLPPSAVQAVLAELPALVNESD MHVAQLAVDFLATVTQAQPASLVEVSGPVLSEL LRLLRSPLLPAGVLAAAEGFLQALVGTRPPCVDY AKLISLLTAPVYEQAVDGGPGLHKQVFHSLARC VAALSAACPQ\EAESTASRLVCDARSPHSSTGVK **VLAFLSLAEVGQVAGPGHERELKAVLLEALGSPS**

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SEQ ID NO:	Method	Predicted beginning nucleotide location corresponding to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \text{\tex{\tex
				EDVRAAASYALGRVGAGSLPDFLPFLLEQIEAEP RRQYLLLHSLKEALGAAQPDSLKPYAEDIWALL FQRCEGAEEGTRGVVAECIGKLVLVNPSFLLPRL RKQLAAGRPHTRSTVITAVKFLISDQPHPIDPLLK SFIAVHNKPSLVRDLLDDILPLLYQETKIRRDLIRE VEMGPFKHTVDDGLDVRKAAFECMYSLLESCLG QLDICEFLNHVEDGLKDHYDIRMLTFIMVARLAT LCPAPVLQRVDRLIEPLRATCTAKVKAGSVKQEF EKQDELKRSAMRAVAALLTIPEVGKSPIMADFSS
3756	A	112	1361	QIRSNPELAALFESIQKDSTSAPSTDSMELS SLEEQQGRHPSFAPKCASQILGRIMITLITEQLQK QTLDELKCTRFSISLPLPDHADISNCGNSFQLVSE GASWRGLPHCSCAEFQ/DQPQLQLPSLRPEPAPQ TT\HRGNSPKEQPFSQVLRPEPPDPEKLPVPPAPPS KRHCRSLSVPVDLSRWQPVWRPAPSKLWTPIKH RGSGGGGGPQVPHQSPPKRVSSL/SVPPSSQCLFS MCPSSHTLQPSFLQPGPGP\DSSRPCAASPQSGSW ESDAESLSPCPPQRRFSLSPSLGPQASRFLPSARSS PASSPELPWRPRGLRNLPRSRSQPCDLDARKTGV KRRHEEDPRRLRPSLDFDKMNQKPYSGGLCLQE TAREGSSISPPWFMACSPPPLSASCSPTGGSSQVL SESEEEEEGAVRWGRQALSKRTLCQRDFGDLDL NLIEEN
3757	A	413	1	PKPMLQQDFT/SLPDQGLDHIAE/NSYFDARSLCA AELVCKEWQQVTSE*MLWKKLIERMVHAYPLW KGLSEKVW/DQHLFKNRPTDGPPNSFHRSLYPKII QVIETIESNWQCG*HTLQRIQCHSEKSKGVYCLQ YDDEK
3758	A	2	613	FVSGSPWRMDGSTERLEARRPAGRLPWSSRQEM TRRPSLMAGRQHGWSAQQSATVANPVPGANPD LLPHFLGEPEDVYIVKNKPVLLVCKAVPATQIFF KCNGEWVRQVDHVIERSTDGSSGLPTMEVRINV SRQQVEKVFGLEEYWCQCVAWSSSGTTKSQKA YIRIAYLRKNFEQEPLAKEVSLEQGIVLPCRPPEGI PPAE
3759	A	1	561	ADDTLHLWNLRQKRPAILHSLKFCRERVTFCHLP FQSKWLYVGTERGNIHIVNVESFTLSGYVIMWN KAIELSSKSHPGPVVHISDNPMDEGKLLIGFESGT VVLWDLKSKKADYRYTYDEAIHSVAWHHEGKQ FICSHSDGTLTIWNVRSPAKPVQTITPHGKQLKD GKKPEPCKPILKVEFXTTR
3760	A .	1	824	LPACRCGCVAGCPSNHGICRCLRASERQVCVMH LKHLRTLLSPQDGAAKVTCMAWSQNNAKFAVC TVDRVVLLYDEHGERRDKFSTKPADMKYGRKS YMVKGMAFSPDSTKIAIGQTDNIIYVYKIGEDWG DKKVICNKFIQTVKFRPVPGTLG*TNIYQYIYL*IQ PGVAFLTSECDFSYCKDGASWLFMVICCLP*SPA VSFPIGD*\SAVTCLQWPAEYIIVFGLAEGKVRLS NTKTNKSSTIYGTESYVVSLTTNCSGKGILSGHA DGYQR
3761	A .	2253	320	PVIQRCSQPYGFSLLISFFLKCVSETSQQPPSRKVF QLLPSFPTLTRSKSHESQLGNRIDDVSSMRFDLSH GSPQMVRRDIGLSVTHRFSTKSWLSQVCHVCQK SMIFGVKCKHCRLKCHNKCTKEAPACRISFLPLT RLRRTESVPSDINNPVDRAAEPHFGTLPKALTKK

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		corresponding to first amino acid residue of peptide sequence	corresponding to last amino acid residue of peptide sequence	N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion
				EHPPAMNHLDSSSNPSSTTFSTPSSPAPFPTSSNPS SATTPP\NPSP\GQR\DSRFNFPSC/AYFIHHR\Q\QFI FPDISAFAHAAPLPEAADGTRLDDQPKADVLEAH EAEAEEPEAGKSEAEDDEDEVDDLPSSRRPWRG PISRKASQTSVYLQEWDIPFEQVELGEPIGQGRW GRVHRGRWHGEVAIRLLEMDGHNQDHLKLFKK EVMNYRQTRHENVVLFMGACMNPPHLAIITSFC KGRTLHSFVRDPKTSLDINKTRQIAQEIIKGMGYL HAKGIVHKDLKSRNVFYDNG\KVVITDFGLF\GIS GVVP\EGRRENQLKLSHDWLCYLAPEIVREMTPG KDEDQLPFSKAADVYAFGTVWYELQARDWPLK NQAAEASIWQIGSGEGMKRVLTSVSLGKEVSEN LSACWAFDLQERPS\FSLLMDMLEKLPKLNRRLS HPGHF*KSADINSSKVVPRFERFGLGVLESSNPK M
3762	A	2	1578	MAHYITFLCMVLVLLLQNSVLAEDGEVRSSCRT APTDLVFILDGSYSVGPENFEIVKKWLVNITKNF DIGPKFIQVGVVQYSDYPVLEIPLGSYDSGEHLTA AVESILYLGGNTKTGKAIQFALDYLFAKSSRFLT KIAVVLTDGKSQDDVKDAAQAARDSKITLFAIG VGSETEDAELRAIANKPSSTYVFYVEDYIAISKIR EVMKQKLCEESVCPTRIPVAARDERGFDILLGLD VNKKVKKRIQLSPKKIKGYEVTSKVDLSELTSNV FPEGLPPSYVFVSTQRFKVKKIWDLWRILTIDG/* PQIAVTLNGVDKILLFTTTSVINGSQVVTFANPQV KTLFDEGWHQIRLLVTEQDVTLYIDDQQIENKPL HPVLGILINGQTQIGKYSGKEETVQFDVQKLRIY CDPEQNNRETACEIPGFCLNGPSDVGSTPAPCICP PGKPGLQGPKGDPGLPGNPGYPGQPGQDGKPVS TESLVISGISGITGYQGIAGTPGVPGSPGIQGARGL PGYKGEPGRDGDK
3763	A	3	1267	CKVWRNPLNLFRGAEYNRYTWVTGREPLTYYD MNLSAQDHQTFFTCDSDHLRPADAIMQKAWRE RNPQARISAAHEALEINECATAYILLAEEEATTIA EAEKLFKQALKAGDGCYRRSQQLQHHGSQYEA QHSVLYLPLQ\TRHQCLGVHQKKASNVCQKTRE DQGSSENDERFNEGVPPSEYVQYP*KPF\KALLEL QAYADVQAVLAKYDDISLPKSATICYTAALLKA RAVSDKFSPEAASRRGLSTAEMNAVEAIHRAVEF NPHVPKYLLEMKSLILPPEHILKRGDSEAIAYAFF HLAHWKRVEGALNLLHCTWEGTFRMIPYPLEKG HLFYPYPICTETADRELLPSFHEVSVYPKKELPFFI LFTAGLCSFTAMLALLTHQFPELMGVFAKAVSV CLEGGLGEWMGKAKGIKAA
3764	A	25	1032 3456	RSADGLCGNKDRERGNEFTRNQQAAQEVVNPK KKMKKKKYVNSGTVTLLSFAVESECTFLDYIKG GTQINFTVAIDFTASNGNPSQSTSLHYMSPYQLN AYALALTAVGEIIQHYDSDKMFPALGFGAKLPPD GRVSHEFPLNGNQENPSCCGIDGILEAYHRSLRT VQLYGPTNFAPVVTHVARNAAAVQDGSQYSVL LIITDGVISDMAQTKEAIVNG\SKLPMSIIIVGVGQ AEFNAMVELDGDDVRISSRGKLAERDIVQFVPFR DYVDRTGNHVLSMARLARDVLAEIPDQLVSYM KAQGIRPRSPPAAPTHSPSQSPARTPPACPLHTHI LGMMDSPKIGNGLPVIGPGTDIGISSLHMVGYLG

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SEQ ID NO:	Method	Predicted beginning nucleotide location corresponding to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion
				KNFDSAKVPSDEYCPACKEKGKLKALKTYRISFQ ESIFLCEDLQCIYPLGSKSLNNLISPDLEECHTPHK PQKRKSLESSYKDSLLLANSKKTRNYIAIDGGKV LNSKHNGEVYDETSSNLPDSSGQQNPIRTADSLE RNEILEADTVDMATTKDPATVDVSGTGRPSPQN EGCTSKLEMPLESKCTSFPQALCVQWKNAYALC WLDCILSALVHSEELKNTVTGLCSKEESIFWRLL TKYNQANTLLYTSQLSGVKDGDCKKLTSEIFAEI ETCLNEVRDEIFISLQPQLRCTLGDMESPVFAFPL LLKLETHIEKLFLYSFSWDFECSQCGHQYQNRH MKSLVTFTNVIPEWHPLNAAHFGPCNNCNSKSQI RKMVLEKVSPIFMLHFVEGLPQNDLQHYAFHFE GCLYQITSVIQYRANNHFITWILDADGSWLECDD LKGPCSERHKKFEVPASEIHIVIWERKISQVTDKE AACLPLKKTNDQHALSNEKPVSLTSCSVGDAAS AETASVTHPKDISVAPRTLSQDTAVTHGDHLLSG PKGLVDNILPLTLEETIQKTASVSQLNSEAFLLEN KPVAENTGILKTNTLLSQESLMASSVSAPCNEKLI QDQFVDISFPSQVVNTNMQSVQLNTEDTVNTKS VNNTDATGLIQGVKSVEIEKDAQLKQFLTPKTEQ LKPERVTSQVSNLKKKETTADSQTTTSKSLQNQS LKENQKKPFVGSWVKGLISRGASFMPLCVSAHN RNTITDLQPSVKGVNNFGGFKTKGINQKASHVSK KARKSASKPPPISKPPAGPPSSNGTAAHPHAHAA SEVLEKSGSTSCGAQLNHSSYGNGISSANHEDLV EGQIHKLRLKLRKKLKAEKKKLAALMSSPQSRT VRSENLEQVPQDGSPNDCESIEDLLNELPYPIDIA NESACTTVPGVSLYSSQTHEEILAELLSPTPVSTE LSENGEGDFRYLGMGDSHIPPPVPSEFNDVSQNT HLRQDHNYCSPTKKNPCEVQPDSLTNNACVRTL NLESPMKTDIFDEFFSSSALNALANDTLDLPHFDE YLFENY
3766	A		1622	AQQIVYRNVMLENYKNLVSLGYQLTKPDVILRL EKGEEPWLVEREIHQETHPDSETAFEIKSSVSSRSI FKDKQSCDIKMEGMARNDLWYLSLEEVWKCRD QLDKYQENPERHLRQVAFTQKKVLTQERVSESG KYGGNCLLPAQLVLREYFHKRDSHTKSLKHDLV LNGHQDSCASNSNECGQTFCQNIHLIQFARTHTG DKSYKCPDNDNSLTHGSSLGISKGIHREKPYECK ECGKFFSWRSNLTRHQLIHTGEKPYECKECGKSF SRSSHLIGHQKTHTGEEPYECKECGKSFSWFSHL VTHQRTHTGDKLYTCNQCGKSF/VHSSRLIRHQR THTGEKPYECPECGKSFRQSTHLILHQRTHVRVR PYECNECGKSYSQRSHLVVHHRIHTGLKPFECKD CGKCFSRSSHLYSHQRTHTGEKPYECHDCGKSFS QSSALIVHQRIHTGEKPYECCQCGKAFIRKNDLIK HQRIHVGEETYKCNQCGIIFSQNSPFIVHQIAHTG EQFLTCNQCGTALVNTSNLIGYQTNHIRENAY
3767	A	3		AQQIVYRNVMLENYKNLVSLGYQLTKPDVILRL EKGEEPWLVEREIHQETHPDSETAFEIKSSVSSRSI FKDKQSCDIKMEGMARNDLWYLSLEEVWKCRD QLDKYQENPERHLRQVAFTQKKVLTQERVSESG KYGGNCLLPAQLVLREYFHKRDSHTKSLKHDLV LNGHQDSCASNSNECGQTFCQNIHLIQFARTHTG DKSYKCPDNDNSLTHGSSLGISKGIHREKPYECK

SEQ ID NO:	Method	Predicted beginning nucleotide location corresponding to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion
			-	ECGKFFSWRSNLTRHQLIHTGEKPYECKECGKSF SRSSHLIGHQKTHTGEEPYECKECGKSFSWFSHL VTHQRTHTGDKLYTCNQCGKSF/VHSSRLIRHQR THTGEKPYECPECGKSFRQSTHLILHQRTHVRVR PYECNECGKSYSQRSHLVVHHRIHTGLKPFECKD CGKCFSRSSHLYSHQRTHTGEKPYECHDCGKSFS QSSALIVHQRIHTGEKPYECCQCGKAFIRKNDLIK HQRIHVGEETYKCNQCGIIFSQNSPFIVHQIAHTG EQFLTCNQCGTALVNTSNLIGYQTNHIRENAY
3768	A	185	2258	SIIIKMSRKISKESKKVNISSSLESEDISLETTVPTD DISSSEEREGKVRITRQLIERKELLHNIQLLKIELS QKTMMIDNLKVDYLTKIEELEEKLNDALHQKQL LTLRLDNQLAFQQKDASKYQELMKQEMETILLR QKQLEETNLQLREKAGDVRRSLRDFELTEEQYIK LKAFPEDQLSIPEYVSVRFYELVNPLRKEICELQV KKNILAEELSTNKNQLKQLTETYEEDRKNYSEV QIRCQRLALELADTKQLIQQGDYRQENYDKVKS ERDALEQEVIELRRKHEILEASHMIQTKERSELSK EVVTLEQTVTLLQKDKEYLNRQNMELSVRCAHE EDRLERLQAQLEESKKAREEMYEKYVASRDHY KTEYENKLHDELEQIRLKTNQEIDQLRNASREMY ERENRNLREARDNAVAEKERAVMAEKDALEKH DQLLDRYREVLQVLSTESKVTEFLHQSKLKSFESE RVQLLQEETARNLTQCQLECEKYQKKLEVLTKE FYSLQASSEKRITELQAQNSEHQARLDIYEKLEK ELDEIIMQTAEIENEDEAERVLFSYGYGANVPTT AKRRLKQSVHLARRVLQLEKQNSLI/LKRSGTSK GPSNTAFTRSLTEANSLLNQTQQPYRYLIESVRQ RDSKIDSLTESIAQL/ERKDVSNLNKEKSALLQTN GIKMALVDLVDQLLNHP
3769		3	2297	DAAEFRVVADAMKVIGFKPEEIQTVYKILAAILH LGNLKFVVDGDTPLIENGKVVSIIAELLSTKTDM VEKALLYRTVATGRDIIDKQHTEQEASYGRDAF AKAIYERLFCWIVTRINDIIEVKNYDTTIHGKNTV IGVLDIYGFEIFDNNSFEQFCINYCNEKLQQLFIQL VLKQEQEEYQREGIPWKHIDYFNNQIIVDLVEQQ HKGIIAILDDACMNVGKVTDEMFLEALNSKLGK HAHFSSRKLCASDKILEFDRDFRIRHYAGDVVYS VIGFIDKNKDTLFQDFKRLMYNSSNPVLKNMWP EGKLSITEVTKRPLTAATLFKNSMIALVDNLASK EPYYVRCIKPNDKKSPQIFDDERCRHQVEYLGLL ENVRVRRAGFAFRQTYEKFLHRYKMISEFTWPN HDLPSDKEAVKKLIERCGFQDDVAYGKTKIFIRT PRTLFTLEELRAQMLIRIVLFLQKVWRGTLARMR YKRTKAALTIIRYYRRYKVKSYIHEVARRFHGVK TMRDYGKHVKWPSPPKVLRRFEEALQTIFNRWR ASQLIKSIPASDLPQVRAKVAAVEMLKGQRADL GLQRAWEGNYLASKPDTPQTSGTFVPVANELKR KDKYMNVLFSCHVRKVNRFSKVEDRAIFVTDRH LYKMDPTKQYKVMKTIPLYNLTGLSVSNGKDQL VVFHTKDNKDLIVCLFSKQPTHESRIGEL\VGVLV NHFKSEKRHLQV\NVTNPVQCSLHGKKCTVSVE TRLNQPQPDFTKNRSGFILSVPGN
3770	A	3	6276	HKVAAPDVVVPTLDTVRHEALLYTWLAEHKPL VLCGPPGSGKTMTLFSALRALPDMEVVGLNFSS

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Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, SEQ ID Method Predicted Predicted end E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine. nucleotide NO: beginning nucleotide location I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, location corresponding T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, corresponding to last amino acid residue of X=Unknown, *=Stop codon, /=possible nucleotide deletion, to first amino acid residue of peptide \=possible nucleotide insertion peptide sequence sequence ATTPELLLKTFDHYCEYRRTPNGVVLAPVQLGK WLVLFCDEINLPDMDKYGTQRVISFIRQMVEHG GFYRTSDOTWVKLERIQFVGACNPPTDPGRKPLS HRFLRHVPVVYVDYPGPASLTQIYGTFNRAMLR LIPSLRTYAEPLTAAMVEFYTMSQERFTQDTQPH YIYSPREMTRWVRGIFEALRPLETLPVEGLIRIWA HEALRLFQDRLVEDEERRWTDENIDTVALKHFP NIDREKAMSRPILYSNWLSKDYIPVDQEELRDYV KARLKVFYEEELDVPLVLFNEVLDHVLRIDRIFR OPOGHLLLIGVSGAGKTTLSRFVAWMNGLSVYQ IKVHRKYTGEDFDEDLRTVLRRSGCKNEKIAFIM DESNVLDSGFLERMNTLLANGEVPGLFEGDEYA TLMTQCKEGAQKEGLMLDSHEELYKWFTSQVIR NLHVVFTMNPSSEGLKDRAATSPALFNRCVLNW FGDWSTEALYQVGKEFTSKMDLEKPNYIVPDYM PVVYDKLPQPPSHREAIVNSCVFVHQTLHQANA RLAKRGGRTMAITPRHYLDFINHYANLFHEKRSE LEEQQMHLNVGLRKIKETVDQVEELRRDLRIKS **OELEVKNAAANDKLKKMVKDQQEAEKKKVMS** QEIQEQLHKQQEVIADKQMSVKEDLDKVEPAVI EAONAVKSIKKOHLVEVRSMANPPAAVKLALES **ICLLLGESTTDWKQIRSIIMRENFIPTIVNFSAEEIS** DAIREKMKKNYMSNPSYNYEIVNRASLACGPMV KWAIAQLNYADMLKRVEPLRNELQKLEDDAKD NQQKANEVEQMIRDLEASIARYKEEYAVLISEAQ AIKADLAAVEAKVNRSTALLKSLSAERERWEKT SETFKNQMSTIAGDCLLSAAFIAYAGYFDQQMR QNLFTTWSHHLQQANIQFRTDIARTEYLSNADER LRWQASSLPADDLCTENAIMLKRFNRYPLIIDPS GQATEFIMNEYKDRKITRTSFLDDAFRKNLESAL RFGNPLLVQDVESYDPVLNPVLNREVRRTGGRV LITLGDQDIDLSPSFVIFLSTRDPTVEFPPDLCSRV TFVNFTVTRSSLQSQCLNEVLKAERPDVDEKRSD LLKLQGEFQLRLRQLEKSLLQALNEVKGRILDDD TIITTLENLKREAAEVTRKVEETDIVMQEVETVS QQYLPLSTACSSIYFTMESLKQIHFLYQYSLQFFL DIYHNVLYENPNLKGVTDHTQRLSIITKDLFQVA FNRVARGMLHQDHITFAMLLARIKLKGTVGEPT YDAEFOHFLRGNEIVLSAGSTPRIQGLTVEQAEA VVRLSCLPAFKDLIAKVQADEQFGIWLDSSSPEQ TVPYLWSEETPATPIGQAIHRLLLIQAFRPDRLLA MAHMFVSTNLGESFMSIMEQPLDLTQIVGTEVKP NTPVLMCSVPGYDASGHVEDLAAEONTQITSIAI GSAEGFNQADKAINTAVKSGRWVMLKNVHLAP **GWLMQLEKKLHSLQPHACFRLFLTMEINPKVPV** NLLRAGRIFVFEPPPGVKANMLRTFSSIPVSRICK SPNERARLYFLLAWFHAIIQERLRYAPLGWSKKY **EFGESDLRSACDTVDTWLDDTAKGRQNISPDKIP** WSALKTLMAQSIYGGRVDNEFDQRLLNTFLERL FTTRSFDSEFKLACKVDGHKDIQMPDGIRREEFV QWVELLPDTQTPSWLGLPNNAERVLLTTQGVD MISKMLKMOMLEDEDDLAYAETEKKTRTDSTS DGRP\AWMRTLHTTASNWLHLIPQTLSHLKRTVE NIKDPLFRFFE\REVKMGAKLLQ\DVRQDLADV\V QVCEGKKKQTNYLRTLI\NELV\KGILP\RSWSHY

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SEQ ID NO:	Method	Predicted beginning nucleotide location corresponding to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion
	·	-		TVPAG\MTVIQWGVPISARRI\KQLQNISL\AAASG GAKELKNIHVCLGGLFVPEAYITATRQYVAQAN SWSLEELCLEVNVTTSQGATLDACSFGVTGLKL QGATCNNNKLSLSNAISTALPLTQLRWVKQTNT EKKASVVTLPVYLNFTRADLIFTVDFEIATKEDPR SFYERGVAVLCTE
3771	A			LPLLHAGFNRRFMENSSIIACYNELIQIEHGEVRS QFKLRACNSVFTALDHCHEAIEITSDDHVIQYVN PAFERMMGYHKGELLGKELADLPKSDKNRADL LDTINTCIKKGKEWQGVYYARRKSGDSIQQHVKI TPVIGQGGKIRHFVSLKKLCCTTDNNKQIHKIHR DSGDNSQTEPHSFRYKNRRKESIDVKSISSRGSDA PSLQNRRYPSMARIHSMTIEAPITKVINIINAAQEN SPVTVAEALDRVLEILRTTELYSPQLGTKDEDPH TSDLVGGLMTDGLRRLSGNEYVFTKNVHQSHSH LAMPITINDVPPCISQLLDNEESWDFNIFELEAITH KRPLVYLGLKVFSRFGVCEFLNCSETTLRAWFQ VIEANYHSSNAYHNSTHAADVLHATAFFLGKER VKGSLDQLDEVAALIAATVHDVDHPGRTNSFL\C NAGSELAVLYNDT\AV\LESHHTALAFQ\LTVKDT K\CNIFKNID/RGNHYRTLRQAIIDMVLATEMTKH FEHVNKFVNSINKPMAAEIEGSDCECNPAGKNFP ENQILIKRMMIKCADVANPCRPLDLCIEWAGRIS EEYFAQTDEEKRQGLPVVMPVFDRNTCSIPKSQI SFIDYFITDMFDAWDAFAHLPALMQHLADNYKH WKTLDDLKCKSLRLPSDRLKPSHRGGLLTDKGH CESQ
3772	A	1013	50	TLVHADGFPSLHITETCLAYREKRIGIDLVHDTVE HELIKEAEIIQGIMALLTRTLEEASEQIRMNRSAK YNLEKDLKDKFVALTIDDICFSLNNNSPNIRYSEN AVRIEPNSVSLEDWLDFSSTNVEKADKQRNNSL MLKALVD\RILSQTANYLRKQCDVVHTAFKNGL KDTKDARDQLADHLAK\VMEEIASQEKNITALEK AILDQEGPAKVAHTRLETRTHRPNVELCRDVAQ YRLMKEVQEITHNVARLKETLA\QAQAELKGLH RRQLALQEEIQVKENTIYIDEVLCMQMRKSIPLR DGEDHGVWAGGLRPDAVC
3773	A .	1	955	AAARESERQLRLRLCVLNEILGTERDYVGTLRFL QSAFLHRIRQNVADSVEKGLTEENVKVLFSNIEDI LEVHKDFLAALEYCLHPEPQSQHELGNVFLKFK DKFCVYEEYCSNHEKALRLLVELNKIPTVRAFLL SCMLLGGRKTTDIPLEGYL\LSPIQRICKYPLLLKE LAKRTPGKHPDHPAVQ\SALQAMKTVCSNINETK RQMEKLEALEAAA/QSHIEGWEGSNLTDICTQLL LQGTLLKISAGNIQERAFFLFDNLLVYCKRKSRV TGSKKSTKRTKSINGSLYIFRGRINTEVMEVENVE DGTGSPSPSLA
3774	A	4254	2061	ELQGDFSVPDVPKSMAWCENSICVGFKRDYYLI RVDGKGSIKELFPTGKQLEPLVAPLADGKVAVG QDDLTVVLNEEGICTQKCALNWTDIPVAMEHQP PYIIAVLPRYVEIRTFEPRLLVQSIELQRPRFITSGG SNIIYVASNHFVWRLIPVPMATQIQQLLQDKQFE LALQLAEMKDDSDSEKQQQIHHIKNLYAFNLFC QKRFDESMQVFAKLGTDPTHVMGLYPDLLPTDY RKQLQYPNPLPVLSGAELEKAHLALIDYLTQKRS

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SEQ ID	Method	Predicted	Predicted end	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid,
NO:		beginning nucleotide	nucleotide location	E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine,
	1	location	corresponding	N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine,
		corresponding	to last amino	T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine,
		to first amino acid residue of	acid residue of peptide	X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion
		peptide sequence	sequence	(~possible indeedude insertion
				QLVKKLNDSDHQSSTSPLMEGTPTIKSKKKLLQII DTTLLKCYLHTNVALVAPLLRLENNHCHIEESEH
			ĺ	VLKKAHKYSELIILYEKKGLHEKALQVLVDQSK
		· · · · · ·		KANSPLKGHERTVQYLQHLGTENLHLIFSYSVW
			1	VLRDFPEDGLKIFTEDLPEVESLPRDRVLGFLIEN
		Ì	1	FKGLAIPYLEHIIHVWEETGSRFHNCLIQLYCEKV
				QGLMKEYLLSFPAGKTPVPAGEEEGELGEYRQK
			1	LLMFLEISSYYDPGRLICDFPFDGLLEERALLLGR
		1		MGKHEQALFIYVHILKDTRMAEEYCHKHYDRN
				KDGNKDVYLSLLRMYLSPPSIHCLGPIKLELLEPK
			(ANLQAALQVLELHHSKLDTTKALNLLPANTQIN
J				DIRIFLEKVLEENAQKKRFNQVLKNLLHAEFLRV\ QEERILHQQVKCIITEEKVCMVCKKKIGNSAFAR
				YPNGVVHYFCS\KEVNPADT
3775	A	1832	839	MSRARGALCRACLALAAALAALLLLPLPLPRAP
			[APARTPAPAPRAPPSRPAAPSLRPDDVFIAVKTTR
				KNHGPRLRLLLRTW\ISRARQQTFIFTDGDDPELE
				LQGGDRVINTNCSAVRTRQALCCKMSVEYDKFI
				ESGRKWFCHVDDDNYVNARSLLHLLSSFSPSQD
				VYLGRPSLDHPIEATERVQGGRTVTTVKFWFAT
				GGAGFCLSRGLALKMSPWASLGSFMSTAEQVRL
				PDDCTVGYIVEGLLGARLLHSPLFHSHLENLQRL PPDTLLQQVTLSHGGPENPQNVVNVAGGFSLHQ
				DPTRFKSIHCLLYPDTDWCPRQKQGAPTSR
3776	A	3	796	PRAKLGTRARNMAGQDAGCGRGGDDYSEDEGD
				SSVSRAAVEVFGKLKDLNCPFLEGLYITEPKTIQE
				LLCSPSEYRLEILEWMCTRVWPSLQDRFSSLKGV
	ł			PTEVKIQEMTKLGHELMLCAPDDQELLKGCACA
	}			QKQLHFMDQLLDTIRSLTIGCSSCSSLMEHFEDT
				REKNEALLGELFSSPHLQMLLNPECDPWPLDMQ PLLNKQSDDWQWASASAKSEEEEKLAELARQLQ
				ESAAKLHALRTEYFAQHEQGAAAGAA\TSAP
3777	A	3	413	SEEDVIEGKTAVIEKRRKKRSSAGVVED/IGGEVQ
				NMLEGVGVDINKALLAKRKRLEMYTKASLRTSN
				QKIEHVWKTQQDQRQKLNQEYSQQFLTLFQQW
				DLDMQKAEEQEEKILVGIMIRFIINQVSSRNGQPS
2770	<u> </u>	120	700	LLL CONTROL AND ADDRESS OF THE PARTY OF THE
3778	A	132	788	SRLPPPPPHLADGRAGARVPRSARLSRWWVQD
				WTHGPIVRPPAAARTMWVNPEEVLLANALWITE RANPYFILQRRKGHAGDGGGGGGLAGLLVGTLD
				VVLDSSARVAPYRILYQTPDSLVYWTIACG\GSR
		1 1		KEITEHWEWLEQNLLQTLSIFENENDITTFVRGKI
	}			QGILAEYNKINDVKEDDDTEKFKEAIVKFHRLFG
				MPEEEKLVNYYSCSYWKG
3779	Α	2	934	CKSCTLFPQNPNLPPPSTRERPPGCKTVFVGGLPE
	1			NATEEIIQEVFEQCGDITAIRKSKKNFCHIRFAEEF
				MVDKAIYLSGYRMRLGSSTDKKDSGRLHVDFA
	1	1		QARDDFYEWECKQRMRAREERHRRKLEEDRLR
	1	1		PPSPPAIMHYSEHEAALLAEKLKDDSKFSEAM\Q
				VLLSWIERGEVNRR\SANQFYSMVQSANSHVRRL MNEKATHEQEMEEAKENFKNALTGILTQFEQIV
	1	1 1	ł	AVFNASTRQKAWDHFSKAQRKNIDIWAK\HSEE
			ľ	LRNAQSEQLMGIRREEEMEMSDDENCDSPTKKM
				RVDESALGAP
3780	A	1	2535	AAQAEREELAAGRMPGGGPQGAPAAAGGGGVS

		r-3	1 80 30 30 30	(A 11-1-1-1-1-1-1-1-1-1-1-1-1-1-1-1-1-1-1
SEQ ID NO:	Method	Predicted beginning nucleotide location corresponding to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \
		sequence		HRAGSRDCLPPAACFRRRRLARRPGYMRSSTGP GIGFLSPAVGTLFRFPGGVSGEESHHSESRARQC GLDSRGLLVRSPVSKSAAAPTVTSVRGTSAHFGI QLRGGTRLPDRLSWPCGPGSAGWQQEFAAMDS SETLDASWEAACSDGARRVRAAGSLPSAELSSNS CSPGCGPEVPPTPPGSHSAFTSSFSFIRLSLGSAGE RGEAEGCPPSREAESHCQSPQEMGAKAASLDGP HEDPRCLSQPFSLLATRVSADLAQAARNSSRPER DMHSLPDMDPGSSSSLDPSLAGCGGDGSSGSGD AHSWDTLLRKWEPVLRDCLLRNRRQMEVISLRL KLQKLQEDAVENDDYDKAETLQQRLEDLEQEKI
	·			SLHFQLPSRQPALSSFLGHLAAQVQAALRRGATQ QASGDDTHTPLRMEPRLLEPTAQDSLHVSITRRD WLLQEKQQLQKEIEALQARMFVLEAKDQQLRRE IEEQEQQLQWQGCDLTPLVGQLSLGQLQEVSKA LQDTLASAGQIPFHAEPPETIRSLQERIKSLNLSLK EITTKVCMSEKFCSTLRKKVNDIETQLPALLEAK MHAISGNHFWTAKDLTEEIRSLTSDREGLEGLLS KLLVLSSRNVKKLGSVKEDYNRLRREVEHQETA YETSVKENTMKYMETLKNKLCSCKCPLLGKVW EADLEACRLLIQCLQLQEARGSLSVEDERQMDD LEGAAPPIPPRLHSEDKRKTPLKESYILSAELGEK CEDIGKKLLYLEDQLHTAIHSHDEDLIQSLRRELQ MVKETLQAMILQLQPAKEAGEREAAASCMTAG VHEAQA
3781	A	3	995	GRRRAGPAHSARMYNMMETELKPPGPQQTSGG GGGNSTAAAAGGNQKNSPDRVKRPMNAFMVW SRGQRRKMAQENPKMHNSEISKRLGAEWKLLSE TEKRPFIDEAKRLRALHMKEHPDYKYRPRRKTK TLMKKDKYTLPGGLLAPGGNSMASGVGVGAGL GAGVNQRMDSYAHMNGWSNGSYSMMQDQLG YPQHPGLNAHGAAQMQPMHRYDVSALQYNSM TSSQTYMNG/SRPTYSMSYSQQGTPGMAPGS\MG SVVKSEASSSPPVVTSSSHSRAPCQAGDLRDMIS MYLPGAEVPEPAAPSRLHMSQHYQSGPVPGTAI NGTLPLSHM
3782		1		FRVPDSCPVVLHSFTQLDPDLPRPESSTQEIGEELI NGVIYSISLRKVQLHHGGNKGQRWLGYENESAL NLYETCKVRTVKAGTLEKLVEHLVPAFQGSDLS YVTIFLCTYRAFTTTQQVLDLLFKRYGRCDALTA SSRYGCILPYSDEDGGPQDQLKNAISSILGTWLD QYSEDFCQPPDFPCLKQLVAYVQLNMPGSDLER RAHLLLAQLEHSEPIEAEPEGEEDWALSPVPALK PTPELELALTPARAPSPVPAPAPEPEPAPTPAPGSE LEVAPAPAPELQQAPEPAVGLESAPAPALELEPA PEQDPAPSQTLELEPAPAPVPSLQPSWPSPVVAEN GLSEEKPHLLVFPPDLVAEQFTLMDAELFKKVVP YHCLGSIWSQRDKKGKEHLAPTIRATVTQFNSV ANCVITTCLGNRSTKAPDRARVVEHWIEVAREC RILKNFSSLYAILSALQSNSIHRLKKTWEDVSRDS FRIFQKLSEIFSDENNYSLSRELLIKEGTSKFATLE MNPKRAQKRPKETGIIQGTVPYLGTFLTDLVML DTAMKDYLYGRLINFEKRRKEFEVIAQIKLLQSA CNNYSIAPDEQFGAWFRAVERLSETESYNLSCEL EPPSESASNTLRTKKNTAIVKRWSDRQAPSTELS

11 C (21 C) 2 C (2 C

SEQ ID NO:	Method	Predicted beginning nucleotide location corresponding to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methlonine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion
				TSGSSHSKSCDQLRCGPYLSSGDIADALSVHSAG SSSSDVEEINISFVPESPDGQEKKFWESASQSSPET SGISSASSSTSSSSASTTPVAATRTHKRSVSGLCNS SSALPLYNQQVGDCCIIRVSLDVDNGNMYKSILV TSQDKAPAVIRKAMDKHNLEEEEPEDYELLQILS DDRKLKIPENANVFYAMNSTANYDFVLKKRTFT KGVKVKHGASSTLPRMKQKGLKIAKGIF
3783	A	3	869	RSGQGKVYGLIGRRRFQQMDVLEGLNLLITISGK RNKLRVYYLSWLRNKILHNDPEVEKKQGWTTV GDMEGCGHYRVVKYERIKFLVIALKSSVEVYAW APKPYHKFMAFKSFADLPHRPLLVDLTVEEGQR LKVIYGSSAGFHAVDVDSGNSYDIYIPVHIQSQIT PHAIIFLPNTDGMEMLLCYEDEGVYVNTYGRIIK DVVLQWGEMPTSVAYICSNQIMGWGEKAIEIRS VETGHLDGVFMHKRAQRLKFLCERNDKVFFASV RSGGSSQVYFMTLNRNCIMNW
3784	A	1213	457	LSPRQVDGLAGLQKGLSLSLLYQFLMNGIRLGTY GLAEAGGYLHTAEGTHSPARSAAAGAMAGVMG AYLGSPIYMVKTHLQAQAASEIAVGHQYKHQG MFQALTEIGQKHGLVGLWRGALGGLPRVIVGSS TQLCTFSSTKDLLSQWEIFPPQSWKLALVAAMM SGIAVVLAMAPFDVACTRLYNQPHRCTGQGP\LY RGILDALLQTARTEGIFGMYKGIGASYFRLGPHTI LSLFFWDQLRSLYYTDTK
3785	A	193	813	RRRGRHSLCGGKMLAYCVQDATVVDVEKRRNP SKHYVYIINVTWSDSTSQTIYRRY\SKFFDLQMQL LD\KFPI\ESGQKDPKQRIIPFLPGKILFRRSHIRDV AVKRLKPIDEYCRALVRLPPHISQCDEVFRFFEAR PEDVNPPKEQGPSPPDAVLPYGVNKGKQELKAG PNWPGRTHHVVNCVTQKCLFVFHFKFSSSGNKE SKSL
3786	A	3785	1632	EFVGRAASTTVVTRIAWRMADAGIRRVVPSDLY PLVLGFLRDNQLSEVANKFAKATGATQQDANAS SLLDIYSFWLNRSAKVPERKLQANGPVAKKAKK KASSSDSEDSSEEEEEVQGPPAKKAAVPAKRVGL PPGKAAAKASESSSSEESSDDDDEEDQKKQPVQ KGVKPQAKAGQAPPKKAKSSDSDSDSSSEDEPP KNQKPKITP\VTVKAQTKAPPKPARA\APKIANGK AASSSSSSSSSSSSDDSEEEKAAATPKKTVPKKQV VAKAPVKAATTPTRKSSSSEDSSSDEEEQKKPM KNKPGPYSSVPPPSAPPPKKSLGTQPPKKAVEKQ QPVESSEDSSDESDSSSEEEKKPPTKAVVSKATTK PPPAKKAAESSSDSSDSDSSSEDDEAPSKPAGTTK NSSNKPAVTTKSPAVKPAAAPKQPVGGGQKLLT RKADSSSSEEESSSSEEEKTKKMVATTKPKATAK AALSLPAKQAPQGSRDSSSDSDSSSSEEEEKTSK SAVKKKPQKVAGGAAPSKPASAKKGKAESSNSS SSDDSSEEEEEKLKGKGSPRPQAPKANGTSALTA QNGKAAKNSEEEEEEKKKAAVVVSKSGSLKKR KQNEAAKEAETPQAKKIKLQTPNTFPKRKKGEK RASSPFRRVREEEIEVDSRVADNSFDAKRGAAGD WGERANQVLKFTKGKSFRHEKTKKKRGSYRGG SISVQVNSIKFDSE
3787	Α	3	5078	IPEG/RALSAEHTSSLVPSLHITTLGQEQAILSGAV PASPSTGTADFPSILTFLQPTENHASPSPVPEMPTL

SEQ ID NO:	Method	Predicted beginning nucleotide location corresponding to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion
		peptide sequence	sequence	PAEGSDGSPPATRDLLLSSKVPNLLSTSWTFPRW KKDSVTAILGKNEEANVTIPLQAFPRKEVLSLHT VNGFVSDFSTGSVSSPIITAPRTNPLPSGPPLPSILS IQATQTVFPSLLAFSSTKPEVYAAAVDHSGLPAS APKQVRASPSSMDVYDSLTIGDMKKPATTDVFW SSLSAETGSLSTESIISGLQQQTNYDLNGHTISTTS WETHLAPTAPPNGLTSAADAIKSQDFKDTAGHS VTAEGFSIQDLVLGTSIEQPVQQSDMTMVGSHID LWPTSNNNHSRDFQTAEVAYYSPTTRHSVSHPQ LQLPNQPAHPLLLTSPGPTSTGSLQEMLSDGTDT GSEISSDINSSPERNASTPFQNILGYHSAAESSISTS VFPRTSSRVLRASQHPKKWTADTVSSKVQPTAA AAVTLFLRKSSPPALSAALVAKGTSSSPLAVASG PAKSSSMTTLAKNVTNKAASGPKRTPGAVHTAF PFTPTYMYARTGHTTSTHTA/IARKHGHCLWPVV YNLP/PP/GKPQAMHTGLPNPTNLEMPRASTPRPL TVTAALTSITASVKATRLPPLRAENTDAVLPAAS AAVVTTGKMASNLECQMSSKLLVKTVLFLTQRR VQISESLKFSIAKGLTQALRKAFHQNDVSAHVDI LEYSHNVTVGYYATKGKLVYLPAVVIEMLGVY GVSNVTADLKQHTPHLQSVAVLASPWNPQPAG YFQLKTVLQFVSQADNIQSCKFAQTMEQRLQKA FQDAERKVLNTKSNLTIQIVSTSNASQAVTLVYV VGRQSTFLNGTVASSLLSQLSAELVGFYLTYPPL TIAEPLEYPNLDISETTRDYWVITVLQGVDNSLV GLHNQSFARVMEQRLAQLFMMSQQQGRRFKRA TTLGSYTVQMVKMQRVPGPKDPAELTYYTLYN GKPLLGTAAAKILSTIDSQRMALTLHHVVLLQAD PVVKNPPNNLWIIAAVLAPIAVVTVIIIITAVLCR KNKNDFKPDTMINLPQRAKPVQGFDYAKQHLG QQGADEEVIPVTQETVVLPLPRDAPQERDVAQD GSTIKTAKSTETRKSRSPSENGSVISNESGKPSSGR RSPQNVMAQQKVTKEEARKRNVPASDEEEGAV LFDNSSKVAAEPFDTSSGSVQLIAIKPTALPMVPP TSDRSQESSAVLNGEVNKALKQKSDIEHYRNKL RLKAKRGYYDFPAVETSKGLTERKKMYEKAP KEMEHYLDPDSELCAPFTESKNRQQMKNSVYRS RQSLNSPSPGETEMDLLVTRERPRRGIRNSGYDT EPEIIEETNIDRVPEPRGYSRSRQVKGHSETSTLSS QPSDDEVRQQMHMLLEEAFSLASAGHAGQSRHQ EAYGSAQHLPYSEVVTSAPGTMTRPRAGVQWVP TYRPEMYQYSLPRPAYRFSQLPEMVMGSPPPPVP PRTGPVAVASLRRSTSDIGSKTRMAESTGPEPAQ LHDSASFTOMSRGPVSVTOLDOSALNYSGNTVP
				AVFAIPAANRPGFTGYFIPTPPSSYRNQAWMSYA GENELPSQWADSVPLPGYIEAYPRSRYPQSSPSRL PRQYSQPANLHPSLEQAPAPSTAASQQSLAENDP SDAPLTNISTAALVKAIREEVAKLAKKQTDMFEF QV
3788	A	2	1737	MKGLYTDAEMKSDNVKDKDAKISFLQKAIDVV VMVSGEPLLAKPARIVAGHEPERTNELLQIIGKC CLNKLSSDDAVRRVLAGEKGEVKGRASLTSRSQ ELDNKNVREEESRVHKNTEDRGDAEIKERSTSRD RKQKEELKEDRMPREKDKDKEKAKENGGNRHR EGERERAKARARPDNERQKDRGNRERDRDSERK

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SEQ ID NO:	Method	Predicted beginning nucleotide location corresponding to first amino acid residue of peptide sequence	Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence	Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion
				KETERKSEGGKEKERLRDRDRERDRDKGKDRDR RRVKNGEHSWDLDRENNREHDKPEKKSASSGE MSKKLSDGTFKDSKAETETEISTRASKSLTTKTS KRRSKNSVEGDSTSDAEGDAGPAGQDKSEVPET PEIPNELSSNIRRIPRPGSARPAPPRVKRQDSMEAL QMDRSGSGKTVSNVITESHNSDNEEDDQFVVEA APQLSEMSEIEMVTAVELEEEEKHGGLVKKILET KKDYEKLQQSPKPGEKERSLFESAWKKEKDIVS KEIEKLRTSIQTLCKSALPLGKIMDYIQEDVDAM QNELQM\YHSENRQHAEALQQEQRITDCAVEP\L KAELA\ELEQLIKD\Q\QDKICAVKANILKNEEKIQ KMVYSINLTSRR
3789	A		4369	MRTLGTCLATLAGLLLTAAGETFSGGCLFDEPYS TCGYSQSEGDDFNWEQVNTLTKPTSDPWMPSGS FMLVNASGRPEGQRAHLLLPQLKENDTHCIDFH YFVSSKSNSPPGLLNVYVKVNNGPLGNPIWNISG DPTRTWNRAELAISTFWPNFYQVIFEVITSGHQG YLAIDEVKVLGHPCTRTPHFLRIQNVEVNAGQFA TFQCSAIGRTVAGDRLWLQGIDVRDAPLKEIKVT SSRRFIASFNVVNTTKRDAGKYRCMINTEGGVGI SNYAEL\VVKEPPVPIAPPQLASVGATYLWIQLN ANSINGDGPIVAREVEYCTASGSWNDRQPVDSTS YKIGHLDPDTEYEISVLLTRPGEGGTGSPGPALRT RTKCADPMRGPRKLEVVEVKSRQITIRWEPFGY NVTRCHSYNLTVHYCYQVGGQEQVREEVSWDT ENSHPQHTITNLSPYTNVSVKLILMNPEGRESQ ELIVQTDEDLPGAVPTESIQGSTFEEKIFLQWREP TQTYGVITLYEITYKAVSSFDPEIDLSNQSGRVSK LGNETHFLFFGLYPGTTYSFTIRASTAKGFGPPAT NQFTTKISAPSMPAYELETPLNQTDNTVTVMLKP AHSRGAPVSVYQIVVEEERPRRTKKTTEILKCYP VPIHFQNASLLNSQYYFAAEFPADSLQAAQPFTIG DNKTYNGYWNTPLLPYKSYRIYFQAASRANGET KIDCVQVATKGAATPKPVPEPEKQTDHTVKIAG VIAGILLFVIIFLGVVLVMKKRKL\AKKKETMSS TRQEIDLWIGELNGPRSYAEQGTKLATRAFSFMD THNLNGRSVSSPSSFTMKTNTLSTSVPNSYYPDE THTMASDTSSLVQSHTYKKREPADVPYQTGQLH PAIRVADLLQHITQMKCAEGYGFKEEYESFFEGQ SAPWDSAKKDENRMKNRYGNIIAYDHSRVRLQT IEGDTNSDYINGNYIDGYHRPNHYIATQGPMQET IYDFWRMVWHENTASIIMVTNLVEVGRVKCCK YWPDDTEIYKDIKVTLIETELLAEYVIRTFAVEKR GVHEIREIRQFHFTGWPDHGVPYHATGLLGFVR QVKSKSPPSAGPLVVHCSAGAGRTGCFIVIDIML DMAEREGVVDIYNCVRELRSRRVNMVQTEEQY VFIHDAILEACLCGDTSVPASQVRSLYYDMNKLD PQTNSSQIKEEFRTLNMVTPTLRVEDCSIALLPRN HEKNRCMDILPPDRCLPFLITIDGESSNYINAALM DSYKQPSAFIVTQHPLPNTVKDFWRLVLDYHCTS VVMLNDVDPAQLCPQYWPENGVHRHGPIQVEF VSADLEEDIISRIFRIYNAARPQDGYRMVQQFQFL GWPMYRDTPVSKRSFLKLIRQVDKWQEEYNGG EGRTVVHCLNGGGRSGTFCAISIVCEMLRHQRTV DVFHAVKTLRNNKPNMVDLLDQYKFCYEVALE

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SEQ ID Method Predicted end nucleotide location corresponding to last amino Method Predicted end nucleotide elocation corresponding to last amino Amino acid sequence (A=Alanine C=Cys E=Glutamic Acid, F=Phenylalanine, G=Cys I=Isoleucine, K=Lysine, L=Leucine, M=Isoleucine, K=Lysine, L=Leucine, M=IT-Threonine, V=Valine, W=Tryptophan	Glycine, H=Histidine, Methionine,
	, R=Arginine, S=Serine,
to first amino acid residue of acid residue of peptide sequence X=Unknown, *=Stop codon, /=possible nucleotide insertion	i, Y=1 yrosine, ucleotide deletion,
sequence YLNSG	
3790 A 261 485 EEQTPLHIASRLGKTEIVQLLLG	OHMAHPDAATTN
GYTPLHISAREGQV\DV\ASVL	
3791 A 1 5874 LPPVTMSGKYIMEEHDSYSDQ	VWSIDELPSKOG
YYLQGNYLRCVAEVGSFEHNI	
KVFMKEVNEVIQKVSGGEQPI	PLWNEHDGTADG
DKPKILLYSLNLQFKGIQVTAT	
LIELELSNRLQTKASPGSSSYLI ALGQIVKHQVYEEAGSDFHQV	
LREEISGSSDREAVLITLNRPIV	
FWLNYK\AAYDNWNEQRMAI	
DMLPGIQQTSAQAFGTPFLQL	TVNDLGICLPITNT
AQSNHTGDLDTGSALVLTIES	
GHFKNFCIRFADGFETSWDDW	
CVVPDGTYEVCSRTTGQAAAI WKMCGIDVHMDPNIGKRLNA	
IDDIADLNSVNIADLSDEDEVE	
RRQAASASQPGELRGRKIMKR	
IDDLKKLGASEGTINQEIQRYC	
DVRKKLRRSSMRAASLKDKW	
ISASGRPPLKRMERASSRVGET GPRVTFNIQDTFPEETELDLLS	
GSCSVFSSPKTPGGFSPGIPFQT	
EDSEKDEKDEDHERERFYIYR	
AAVHQLFTERWPTTPVNRSLS	
IRVEIDSGKCVLHPTTLLQEHD	
SLDQDSPSKKKKFQTNYASTT QTKPSDLETTVFYIPGVDVKLI	
ASRGSSLPRTLSKESKLYGMK	
VQSKTNTLLPPQPPPIPAAKGK	
WVALQSLPEEMVISPCLLDFLI	
NYTAVSSQDEDMGHFEIPDPM	
AYSSFPVDVVVYVRVQPSQIK KLPSLDLVFSSNRGELETLGTI	
TQSGTKTSASKTGIPGSSGLGS	
LTSSSSSGLSFTACMSDFSL	YVFHPYGAGKQIT
AVSGLTPGSGGLGNVDEEPTS	
FVKVSLSRIRRSGGASFFESQS NISAVCDIGSASFKYDMRRLSI	
ARRLFLGDQTINLPTSGPGTPD	
RKAYCKTWEQPSQSASFTHM	
STMSPGTVGQSLKSPASIRSRS	VSDSSVPRRDSLS
KTSTPFNKSNKAASQQGTPWE	
NVQMNMSNVMGNTTWTTSG	
DREISMSVGLGRSQLDSKGGV VAHISEHPNQQPSHKIQITMGS	
MGIFSNADLKLQDEWKVNLY	
VHGDLKWDIFQVMISRSTTPD	
QQFDTSKRALSTWGPVPYLPP	KTMTSNLEKSSQE
QLLDAAHHRHWPGVLKVVSC	
MQFGGSMSLHGNHMTLACFH	
HLEEPNIAFWTEAQKIWEDGS HLGHNTMVTKPCGALESPMA	
HGVASVKEWFNYVTATRNEE	

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| SEQ ID<br>NO: | Method | Predicted beginning nucleotide location corresponding to first amino acid residue of peptide sequence | Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence | Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \text{\tex{\tex |
|---------------|--------|-------------------------------------------------------------------------------------------------------|------------------------------------------------------------------------------------------------|--------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
|               |        |                                                                                                       | i.                                                                                             | ENSTTVKNSSLLSGFRGGSSYNHETETIFALPRM QLDFKSIHVQEPQEPSLQDASLKPKVECSVVTEF TDHICVTMDAELIMFLHDLVSAYLKEKEKAIFPP RILSTRPGQKSPIIIHDDNSSDKDREDSITYTTVDW RDFMCNTWHLEPTLRLISWTGRKIDPVGVDYILQ KLGFHHARTTIPKWLQRGVMDPLDKVLSVLIKK LGTALQDEKEKKGKDKEEH                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       |
| 3792          | Α .    | 1                                                                                                     | 364                                                                                            | QNGSTPLHHAASKNRHEIALMLLEGGANPDGKD<br>HYEATAKHQATAKGNFKMIHILLYYKASTIIQDT<br>EGNTPPHLVCD\RVEEAKLLVSQGA\SIYIENKEE<br>KDP/LQVAKGALGLVLKRMVEG                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     |
| 3793          | Ā      | 2                                                                                                     | 340                                                                                            | DIVPNPKMAPLGDEAPTLEKVLTPELSEEEVSTR<br>DDIQFHHFSSEEALQKVKYFVAKEDPSSQEEAHT<br>PEAPPPQPPSSERCLGEMKCTLVRGDSSPRQAEL<br>KSGPASRPAL                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 |
| 3794          | A      | 421                                                                                                   | 158                                                                                            | SYWVGEDYTYKFFEVILIDPFHKAIRRNPDTQWI<br>SKAVYKHREMCGLTSTGRKSHGLEKDRMFPHAI<br>GGSCRAA*RRRKTLQFPCYH                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              |
| 3795          | A      | 24                                                                                                    | 592                                                                                            | GGMDSRVSGTTSNGETKPVYPVMEKKEEDGTLE<br>RGHWNNKMEFVLSVAGEIIGLGNVWRFPYLCYK<br>NGGGAFFIPYLVFLFTCGIPVFLLETALGQYTSQG<br>GVTAWRKICPIFEGIGYASQMIVILLNVYYIIVLA<br>WALFYLFSSFTIDLPWGGCYHEWNTEHCMEFQK<br>TNGSLNGTSENATSPVIEFW                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            |
| 3796          | Α      | 3                                                                                                     | 592                                                                                            | KPASTYSTSQPSMAPLLPIRTLPLILILLALLSPGA<br>ADFNISSLSGLLSPALTESLLVALPPCHLTGGNAT<br>LMVRRANDSKVVTSSFVVPPCRGRRELVSVVDS<br>GAGFTVTRLSAYQVTNLVPGTKFYISYLVKKGT<br>ATESSREIPMFTLPRRNMESIGLGMARTGGMVVI<br>TVLLSVAMFLLVLGFIIALALGSRK                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     |
| 3797          | A      | 1                                                                                                     | 1556                                                                                           | ATRLLRGSGSWGCSRLRFGPPAYRRFSSGGAYPN IPLSSPLPGVPKPVFATVDGQEKFETKVTTLDNGL RVASQNKFGQFCTVGILINSGSRYEAKYLSGIAH FLEKLAFSSTARFDSKDEILLTLEKHGGICDCQTS RDTTMYAVSADSKGLDTVVALLADVVLQPRLT DEEVEMTRMAVQFELEDLNLRPDPEPLLTEMIHE AAYRENTVGLHRFCPTENVAKINREVLHSYLRN YYTPDRMVLAGVGVEHEHLVDCARKYLLGVQP AWGSAEAVDIDRSVAQYTGGIAKLERDMSNVSL GPTPIPELTHIMVGLESCSFLEEDFIPFAVLNMMM GGGGSFSAGGPGKGMFSRLYLNVLNRHHWMYN ATSYHHSYEDTGLLCIHASADPRQVREMVEIITK EFILMGGTVDTVELERAKTQLTSMLMMNLESRP VIFEDVGRQVLATRSRKLPHELCTLIRNVKPEDV KRVASKMLRGKPAVAALGDLTDLPTYEHIQTAL SSKDGRLPRTYRLFR                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        |
| 3798          |        | 73                                                                                                    | 759                                                                                            | KRLVEAGVPRTFDGIVGEGGAQSRSCWPWGVTA QTPAFSADSLNCLKNCMSITMGSVRPSVEQFHKY LPWFLNDRPNIKCPKGGLAAYSTSVNLTSDGQV LASRFMAYHKPLKNSQDYTEALRAARELAANIT ADLRKVPGTDPAFEVFPYTITNVFYEQYLTILPEG LFMLSLCLVPTFAVSCLLLGLDLRSGLLNLLSIV MILVDTVGFMALWGISYNAVSLINLVS                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  |
| 3799          | A      | 73                                                                                                    | 759                                                                                            | KRLVEAGVPRTFDGIVGEGGAQSRSCWPWGVTA<br>QTPAFSADSLNCLKNCMSITMGSVRPSVEQFHKY                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      |

| SEQ ID<br>NO: | Method | Predicted beginning nucleotide location corresponding to first amino acid residue of peptide sequence | Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence | Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion                                                                                                                               |
|---------------|--------|-------------------------------------------------------------------------------------------------------|------------------------------------------------------------------------------------------------|------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
|               |        |                                                                                                       |                                                                                                | LPWFLNDRPNIKCPKGGLAAYSTSVNLTSDGQV<br>LASRFMAYHKPLKNSQDYTEALRAARELAANIT<br>ADLRKVPGTDPAFEVFPYTITNVFYEQYLTILPEG<br>LFMLSLCLVPTFAVSCLLLGLDLRSGLLNLLSIV<br>MILVDTVGFMALWGISYNAVSLINLVS                                                                                                                                                                                                                                                                                                                         |
| 3800          | A      | 250                                                                                                   | 1032                                                                                           | GIFRSLRVLFPLFSVGRPQFARSLSAAPQLSDTAD TMGFGDLKSPAGLQVLNDYLADKSYIEGYVPSQ ADVAVFEAVSSPPPADLCHALRWYNHIKSYEKE KASLPGVKKALGKYGPADVEDTTGSGATDSKD DDDIDLFGSDDEEESEEAKRLREERLAQYESKKA KKPALVAKSSILLDVKPWDDETDMAKLEECVRS IQADGLVWGSSKLVPVGYGIKKLQIQCVVEDDK VGTDMLEEQITAFEDYVQSMDVAAFNKI                                                                                                                                                                                                                               |
| 3801          | A      | 155                                                                                                   | 656                                                                                            | SREMELVTFRDVAIEFSPEEWKCLDPAQQNLYR<br>DVMLENYRNLVSLGFVISNPDLVTCLEQIKEPCN<br>LKIHETAAKPPAICSPFSQDLSPVQGIEDSFHKLIL<br>KRYEKCGHENLQLRKGCKRVNECKVQKGVNNG<br>VYQCLSTTQSKIFQCNTCVRVFSTSSHSNKHK                                                                                                                                                                                                                                                                                                                    |
| 3802          | A .    |                                                                                                       | 1428                                                                                           | VTVSPETHMDLTKGCVTFEDIAIYFSQDEWGLLD EAQRLLYLEVMLENFALVASLGCGHGTEDEETP SDQNVSVGVSQSKAGSSTQKTQSCEMCVPVLKD ILHLADLPGQKPYLVGECTNHHQHQKHHSAKKS LKRDMDRASYVKCCLFCMSLKPFRKWEVGKDL PAMLRLLRSLVFPGGKKPGTITECGEDIRSQKSH YKSGECGKASRHKHTPVYHPRVYTGKKLYECSK CGKAFRGKYSLVQHQRVHTGERPWECNECGKF FSQTSHLNDHRRIHTGERPYECSECGKLFRQNSS LVDHQKIHTGARPYECSQCGKSFSQKATLVKHQ RVHTGERPYKCGECGNSFSQSAILNQHRRIHTGA KPYECGQCGKSFSQKATLIKHQRVHTGERPYKC GDCGKSFSQSSILIQHRRIHTGARPYECGQCGKSF SQKSGLIQHQVVHTGERPYECNKCGNSFSQCSSL IHHQKCHNT |
| 3803          | A      | 193                                                                                                   | 617                                                                                            | LFPFLGSESKNGEADSSDKEMKHGQKSPTGKQTS<br>QHLKRLKKSGLGHLKWTKAEDIDIETPGSILVNT<br>NLRALINKHTFASLPQHFQQYLLLLLPEVDRQMG<br>SDGILRLSTSALNNEFFAYAAQGWKQRLAEGKF<br>VFSIIM                                                                                                                                                                                                                                                                                                                                              |
| 3804          | A      | 197                                                                                                   | 479                                                                                            | SSSRASPPEHPSSQAHCGPLVLSHACPEVTNKWS<br>TGSSSSPNSSWVSSPLQPEGLSGSSRMKGGSATKI<br>LLETLLLAAHMTADQGIASSQRCLL                                                                                                                                                                                                                                                                                                                                                                                                     |
| 3805          | A      | 1                                                                                                     | 385                                                                                            | QSADTLFPGDINFNVSGLFSAVTLQDTVSDRLAS<br>EELPSTAVPTPATTPAPAPAPAPATAPALVSAAT<br>KERTESEVPPRPASPKVTRSPPETAAPVEDMARR<br>SELAVGGEEGTEGGRGEGTGSPMSSY                                                                                                                                                                                                                                                                                                                                                               |
| 3806          | A .    | 47                                                                                                    | 1033                                                                                           | LQGDTWHLSFLSHFSRLHGGVPGRGLLEGNLLQ PQAPGHDMTSIPFPGDRLLQVDGVILCGLTHKQA VQCLKGPGQVARLVLERRVPRSTQQCPSANDSM GDERTAVSLVTALPGRPSSCVSVTDGPKF*SSN* KRIANGLGFSFVQMEKESCSHLKSDLVRIKRLFP GHPAEENGAIAAGDIILGREWEGPRKASSSRCRG SWAMQLSVQAGPSFASYYPAAVEVLHLLRGAPQ EVTLLLCRPPPGALPELEQEWQTPELSADKEFTR ATCTDSCTSPILGSRGQLGGTVPPQMQGKAWGL RPESSQKAIREGTMGAKTERDLGPVP                                                                                                                                                          |

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| 3807          | A        | 656                                                                                                   | 1238                                                                                           | RCPSLLPPSWPLPTLQTLTRTPGNKAIAGGAGLW<br>AVLWGSERTPPYR*GN*NQRGAVPCLRPHRLRP<br>QDKFLVLASDGLWDMLSNEDVVRLVVGHLAEA<br>DWHKTDLAQRPANLGLMQSLLLQRKASGLHEA<br>DQNAATRLIRHAIGNNEYGEMEAERLAAMLTLP<br>EDLARMYRDDITVTVVYFNSESIGAYYKGG                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   |
| 3808          | <b>A</b> | 26                                                                                                    | 2195                                                                                           | SQYSESVAGRQASPERLLGSYHAMASTVEGGDT ALLPEFPRGPLDAYRARASFSWKELALFTEGEG MLRFKKTIFSALENDPLFARSPGADLSLEKYREL NFLRCKRIFEYDFLSVEDMFKSPLKVPALIQCLG MYDSSLAAKYLLHSLVFGSAVYSSGSERHLTYIQ KIFRMEIFGCFALTELSHGSNTKAIRTTAHYDPAT EEFIIHSPDFEAAKFWVGNMGKTATHAVVFAKL CVPGDQCHGLHPFIVQIRDPKTLLPMPGVMVGDI GKKLGQNGLDNGFAMFHKVRVPRQSLLNRMGD VTPEGTYVSPFKDVRQRFGASLGSLSSGRVSIVSL AILNLKLAVAIALRFSATRRQFGPTEEEEIPVLEY PMQQWRLLPYLAAVYALDHFSKSLFLDLVELQR GLASGDRSARQAELGREIHALASASKPLASWTT QQGIQECREACGGHGYLAMNRLGVLRDDNDPN CTYEGDNNILLQQTSNYLLGLLAHQVHDGACFR SPLKSVDFLDAYPGILDQKFEVSSVADCLDSAVA LAAYKWLVCYLLRETYQKLNQEKRSGSSDFEAR NKCQVSHGRPLALAFVELTVVQRFHEHVHQPSV PPSLRAVLGRLSALYALWSLSRHAALLYRGGYF SGEQAGEVLESAVLALCSQLKDDAVALVDVIAP PDFVLDSPIGRADGELYKNLWGAVLQESKVLER ASWWPEFSVNKPVIGSLKSKL |
| 3809          | A        | 117                                                                                                   | 830                                                                                            | CFGIMERVGCTLTTTYAHPRPTPTNFLPAISTMAS<br>SYRDRFPHSNLTHSLSLPWRPSTYYKVASNSPSV<br>APYCTRSQRVSENTMLPFVSNRTTFFTRYTPDDW<br>YRSNLTNYQESNTSRHNSEKLRVDTSRLIQDKYQ<br>QTRKTQADTTQNLGERVNDIGFWKSEIIHELDEM<br>IGETNALTDVKKRLERALMETEAPLQVARECLF<br>HREKRMGIDLVHDEVEAQLLTVNVGEMHQSQA<br>A                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                |
| 3810          | A        | 3                                                                                                     | 518                                                                                            | VIQELEGGSGADLGEHSCRPASQPRFPRPAEARS HPATRRPASGPAMGKTNSKLAPEVLEDLVQNTE FSEQELKQWYKGFLKDCPSGILNLEEFQQLYIKF FPYGDASKFAQHAFRTFDKNGDGTIDFREFICAL SVTSRGSFEQKLNWAFEMYDLDGDGRITRLEML EIIE                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        |
| 3811          | A        | 81                                                                                                    | 1147                                                                                           | GCGYGCSGAGGAAIGEPMAKWGEGDPRWIVEE RADATNVNNWHWTERDASNWSTDKLKTLFLAV QVQNEEGKCEVTEVSKLDGEASINNRKGKLIFFY EWSVKLNWTGTSKSGVQYKGHVEIPNLSDENSV DEVEISVSLAKDEPDTNLVALMKEEGVKLLREA MGIYISTLKTEFTQGMILPTMNGESVDPVGQPAL KTEERKAKPAPSKTQARPVGVKIPTCKITLKETFL TSPEELYRVFTTQELVQAFTHAPATLEADRGGKF HMVDGNVSGEFTDLVPEKHIVMKWRFKSWPEG HFATITLTFIDKNGETELCMEGRGIPAPEEERTRQ GWQRYYFEGIKQTFGYGARLF                                                                                                                                                                                                                                                                                                                                                                                            |
| 3812          | A        | 20                                                                                                    | 558                                                                                            | PCGTAASTHAYDRRAKCRQQQQQQQQQGQNKV<br>RPAKKKTSPAREVSSESGTSGQFTPPSSTSVPTIAS                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 |

| SEO ID | Method   | Predicted                                                                                   | Predicted end                                                                    | Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid,                                                                                                                                                                                                                                                                                                                                                      |
|--------|----------|---------------------------------------------------------------------------------------------|----------------------------------------------------------------------------------|------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| NO:    | , ACTION | beginning nucleotide location corresponding to first amino acid residue of peptide sequence | nucleotide location corresponding to last amino acid residue of peptide sequence | E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion                                                                                                 |
|        |          |                                                                                             |                                                                                  | SSAPVSIWSPASISPLSDPLSTSSSCMQRSYPMTYT<br>QASGYSQGYAGSTSYFGGMDCGSYLTPMHHQL<br>PGPGATLSPMGTNAVTSHLNQSPASLSTQGYGAS<br>KLWGFNFNH                                                                                                                                                                                                                                                                                      |
| 3813   | A        | 1                                                                                           | 1016                                                                             | CTEPPRRSTRTPAALASLRPYTDYVVVSDQILQES EDFFTLIESHEGKPLKLMVYNSKSDSCREVTVTP NAAWGGEGSLGCGIGYGYLHRIPTQPPSYHKKPP GTPPPSALPLGAPPPDALPPGPTPEDSPSLETGSRQ SDYMEALLQAPGSSMEDPLPGPGSPSHSAPDPDG LPHFMETPLQPPPPVQRVMDPGFLDVSGISLLDN SNASVWPSLPSSTELTTTAVSTSGPEDICSSSSSHE RGGEATWSGSEFEVSFLDSPGAQAQADHLPQLT LPDSLTSAASPEDGLSAELLEAQAEEEPASTEGLD TGTEAEGLDSQAQISTTE*HPGL*QGP                                                      |
| 3814   | A        | 2                                                                                           | 884                                                                              | VFWQVRNAGSSPLSAACPLFRTPAPQPCGSWGR CCIPHASTGCRPMAERGELDLTGAKQNTGVWLV KVPKYLSQQWAKASGRGEVGKLRIAKTQGRTE VSFTLNEDLANIHDIGGKPASVSAPREHPFVLQSV GGQTLTVFTESSSDKLSLEGIVVQRAECRPAASE NYMRLKRLQIEESSKPVRLSQQLDKVVTTNYKP VANHQYNIEYERKKKEDGKRARADKQHVLDML FSAFEKHQYYNLKDLVDITKQPVVYLKEILKEIG VQNVKGIHKNTWELKPEYRHYQGEEKSD                                                                                                   |
| 3815   | Α        | 17                                                                                          | 411                                                                              | NIGDWEDIGKSPERIIQYYGPATWAQDGSRGYCT<br>PIYMLNHIIRLQAVLEIIMNERANALDLLAQQTTK<br>MRNANYQNRLALDYLLAHEGGV*GKFSLTNCC<br>LEIDDNGKAIMEITARMRKLAHIPVQTWER                                                                                                                                                                                                                                                                  |
| 3816   | A        | 3                                                                                           |                                                                                  | SHWQRRDRRCVRNMAERGRKRPCGPGEHGQRI EWRKWKQQKKEEKKKWKDLKLMKKLERQRAQ EEQAKRLEEEEAAAEKEDRGRPYTLSVALPGSIL DNAQSPELRTYLAGQIARACAIFCVDEIVVFDEE GQDAKTVEGEFTGVGKKGQACVQLARILQYLEC PQYLRKAFFPKHQDLQFAGLLNPLDSPHHMRQD EESEFREGVVVDRPTRPGHGSFVNCGMKKEVKI DKNLEPGLRVTVRLNQQQHPDCKTYHGKVVSS QDPRTKAGLYWGYTVRLASCLSAVFAEAPFQDG YDLTIGTSERGSDVASAQLPNFRHALVVFGGLQG LEAGADADPNLEVAEPSVLFDLYVNTCPGQGSR TIRTEEAILISLAALQPGLIQAGARHT |
| 3817   | A        | 246                                                                                         | 1197                                                                             | FLSAGMSNFTHYAYLLMIESLMLGKVPPHVPSH HFIFHDDGSARQKGESDYKVIIQQWFSKSGPWTT SSNVTWGLLELQQSISESAVLTIPPGDSGAGSNLI TMFLRNRKETDLCSGRSKVNRGWNSGRCKQRG KTEQPGEPLEHVYVTIKHAVALESRHQKGELQC LIKMCIPLSKPLQMFFSPPHWEAWLQRVQQLAK NTRYFRQRLQEMGFIIYGNENASVVPLLLYMPG KVAAFARHMLEKKIGVVVVGFPATPLAEARARF CVSAAHTREMLDTVLEALDEMGDLLQLKYSRH KKSARPELYDETSFELED                                                                            |
| 3818   | A        | 215                                                                                         | 789                                                                              | NPQSSSSEGSSEIFQVNGHNRLLVQRSEVTQAPG<br>QYTVDVEGHGCTFIQATLKYNVLLPKKASGFSLS<br>LEIVKNYSSTAFDLTVTLKYTGIRNKSSMVVIDV<br>KMLSGFTPTMSSIEELENKGQVMKTEVKNDHVL<br>FYLENVFGRADSFTFSVEQSNLVFNIQPAPGMVY<br>DYYEKEEYALAFYHINSSSVSE                                                                                                                                                                                              |

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Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, Predicted end SEQ ID Method Predicted NO: beginning nucleotide E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, location nucleotide corresponding N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, location T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, corresponding to last amino X=Unknown, \*=Stop codon, /=possible nucleotide deletion. acid residue of to first amino \=possible nucleotide insertion acid residue of peptide peptide sequence sequence 1483 RIPDSIISRGVQGLPRDTASLSTTPSESPRAQATSR 3819 LSTASCPTPKVQSRCSSKENILRASHSAVDITKVA RRHRMSPFPLTSMDKAFITVLEMTPVLGTEIINYR DGMGRVLAODVYAKDNLPPFPASVKDGYAVRA ADGPGDRFIIGESQAGEQPTQTVMPGQVMRVTT GAPIPCGADAVVQVEDTELIRESDDGTEELEVRIL VOARPGODIRPIGHDIKRGECVLAKGTHMGPSEI GLLATVGVTEVEVNKFPVVAVMSTGNELLNPED DLLPGKIRDSNRSTLLATIQEHGYPTINLGIVGDN PDDLLNALNEGISRADVIITSGGVSMGEKDYLKQ VLDIDLHAOIHFGRVFMKPGLPTTFATLDIDGVR KIIFALPGNPVSAVVTCNLFVVPALRKMQGILDP RPTIIKARLSCDVKLDPRPEYHRCILTWHHQEPLP WAOSTGNOMSSRLMSMRSANGLLMLPPKTEQY VELHKGEVVDVMVIGRL POEPALKSEFSQVASNTIPLPLPQPNTCKDNGPCK 3820 A 2216 487 QVCSTVGGSAICSCFPGYAIMADGVSCEDQDECL **MGAHDCSRRQFCVNTLGSFYCVNHTVLCADGYI** LNAHRKCVDINECVTDLHTCSRGEHCVNTLGSF **HCYKALTCEPGYALKDGECEDVDECAMGTHTC QPGFLCQNTKGSFYCQARQRCMDGFLQDPEGNC VDINECTSLSEPCRPGFSCINTVGSYTCQRNPLIC** ARGYHASDDGTKCVDVNECETGVHRCGEGQVC HNLPGSYRCDCKAGFQRDAFGRGCIDVNECWAS PGRLCQHTCENTLGSYRCSCASGFLLAADGKRC **EDVNECEAQRCSQECANTYGSYQCYCRQGYQLA EDGHTCTDIDECAQGAGILCTFRCLNVPGSYQCA** CPEQGYTMTANGRSCKDVDECALGTHNCSEAET CHNIQGSFRCLRFECPPNYVQVSKTKCERTTCHD FLECQNSPARITHYQLNFQTGLLVPAHIFRIGPAP AFTGDTIALNIIKGNEEGYFGTRRLNAYTGVVYL **ORAVLEPRDFALDVEMKLWRQGSVTTFLAKMHI** FFTTFAL 3821 A 2216 487 PQEPALKSEFSQVASNTIPLPLPQPNTCKDNGPCK QVCSTVGGSAICSCFPGYAIMADGVSCEDQDECL MGAHDCSRRQFCVNTLGSFYCVNHTVLCADGYI LNAHRKCVDINECVTDLHTCSRGEHCVNTLGSF **HCYKALTCEPGYALKDGECEDVDECAMGTHTC** OPGFLCONTKGSFYCOARQRCMDGFLQDPEGNC **VDINECTSLSEPCRPGFSCINTVGSYTCQRNPLIC** ARGYHASDDGTKCVDVNECETGVHRCGEGQVC HNLPGSYRCDCKAGFQRDAFGRGCIDVNECWAS PGRLCOHTCENTLGSYRCSCASGFLLAADGKRC **EDVNECEAQRCSQECANIYGSYQCYCRQGYQLA EDGHTCTDIDECAQGAGILCTFRCLNVPGSYQCA CPEQGYTMTANGRSCKDVDECALGTHNCSEAET** CHNIQGSFRCLRFECPPNYVQVSKTKCERTTCHD FLECONSPARITHYOLNFOTGLLVPAHIFRIGPAP AFTGDTIALNIIKGNEEGYFGTRRLNAYTGVVYL QRAVLEPRDFALDVEMKLWRQGSVTTFLAKMHI **FFTTFAL** 3822 2502 1540 MAAATRGCRPWGSLLGLLGLVSAAAAAWDLAS À LRCTLGAFCECDFRPDLPGLECDLAQHLAGQHL AKALVVKALKAFVRDPAPTKPLVLSLHGWTGTG KSYVSSLLAHYLFQGGLRSPRVHHFSPVLHFPHP

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|---------------|--------|-------------------------------------------------------------------------------------------------------|------------------------------------------------------------------------------------------------|----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
|               |        |                                                                                                       |                                                                                                | SHIERYKKOLKSWVQGNLTACGRSLFLFDEMDK<br>MPPGLMEVLRPFLGSSWVVYGTNYRKAIFIFISN<br>TGGEQINQVALEAWRSRRDREEILLQELEPVISR<br>AVLDNPHHGFSNSGIMEERLLDAVVPFLPLQRHH<br>VRHCVLNELAQLGLEPRDEVVQAVLDSTTFFPE<br>DEQLFSSNGCKTVASRIAFFL                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                |
|               | A      |                                                                                                       | 3174                                                                                           | YGCEKTTEGRIPLKNIYRLFSADRKRVETALEAC SLPSSRNDSIPQEDFTPEVYRVFLNNLCPRPEIDNI FSEFGAKSKPYLTVDQMMDFINLKQRDPRLNEIL YPPLKQEQVQVLIEKYEPNNSLARKGQISVDGFM RYLSGEENGVVSPEKLDLNEDMSQPLSHYFINSS HNTYLTAGQLAGNSSVEMYRQVLLSGCRCVELD CWKGRTAEEEPVITHGFTMTTEISFKEVIEAIAEC AFKTSPFPILLSFENHVDSPKQQAKMAEYCRLIFG DALLMEPLEKYPLESGVPLPSPMDLMYKILVKN KKKSHKSSEGSGKKKLSEQASNTYSDSSSMFEPS SPGAGEADTESDDDDDDDDCKKSSMDEGTAGSE AMATEEMSNLVNYIQPVKFESFEISKKRNKSFEM SSFVETKGLEQLTKSPVEFVEYNKMQLSRIYPKG TRVDSSNYMPQLFWNAGCQMVALNFQTMDLA MQINMGMYEYNGKSGYRLKPEFMRRPDKHFDP FTEGIVDGIVANTLSVKIISGQFLSDKKVGTYVEV DMFGLPVDTRRKAFKTKTSQGNAVNPVWEEEPI VFKKVVLPTLACLRIAVYEEGGKFIGHRILPVQAI RPGYHYICLRNERNQPLTLPAVFVYIEVKDYVPD TYADVIEALSNPIRYVNLMEQRAKQLAALTLEDE EEVKKEADPGETPSEAPSEARTTPAENGVNHTTT LTPKPPSQALHSQPAPGSVKAPAKTEDLIQSVLTE VEAQTIEELKQQKSFVKLQKKHYKEMKDLVKR HHKKTTDLIKEHTTKYNEIQNDYLRRAALEKS AKKDSKKKSEPSSPDHGSSTIEQDLAALDAEMTQ KLIDLKDKQQQQLLNLRQEQYYSEKYQKREHIK LLIQKLTDVAEECQNNQLKKLKEICEKEKKELKK KMDKKRQEKITEAKSKDKSQMEEEKTEMIRSYI QEVVQYIKRLEEAQSKRQEKLVEKHKEIRQQILD EKPKLQVELEQEYQDKFKRLPLEILEFVQEAMKG KISEDSNHGSAPLSLSSDPGKVNHKTPSSEELGGD IPGKEFDTPL |
| 3824          | A      | 1                                                                                                     | 426                                                                                            | ILHWFVHRWSGRNNREKIGVHVGFEEILNMEPY<br>CCRETLKSLRPECFIYDLSAVVMHHGKGFGSGH<br>YTAYCYNSEGGFWVHCNDSKLSMCTMDEVCKA<br>QAYILFYTQRVTENGHSKLLPPELLLGSQHPNED<br>ADTSSNEILS                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   |
| 3825          | A      | 3                                                                                                     | 364                                                                                            | GIRAKFPNKIPVVVERYPRETFLPPLDKTKFLVPQ<br>ELTMTQFLSIIRSRMVLRATEAFYLLVNNKSLVS<br>MSATMAEIYRDYKDEDGFVYMTYASQETFGCLE<br>SAAPRDGSSLEDRPLHPL                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             |
| 3826          | Α      | 1                                                                                                     | 1237                                                                                           | PEKKFERECREAEKAQQSYERLDNDTNATKADV EKAKQQLNLRTHMADENKNEYAAQLQNFNGEQ HKHFYVVIPQIYKQLQEMDERRTIKLSECYRGFA DSERKVIPIISKCLEGMILAAKSVDERRDSQMVV DSFKSGFEPPGDFPFEDYSQHIYRTISDGTISASKQ ESGKMDAKTTVGKAKGKLWLFGKKPKGPALED FSHLPPEQRRKKLQQRIDELNRELQKESDQKDAL NKMKDVYEKNPQMGDPGSLQPKLAETMNNIDR                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               |

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|               |        | ·                                                                                                     |                                                                                                | LRMEIHKNEAWLSEVEGKTGGRGDRRHSSDINH<br>LVTQGRESPEGSYTDDANQEVRGPPQQHGHHNE<br>FDDEFEDDDPLPAIGHCKAIYPFDGHNEGTLAMK<br>EGEVLYIIEEDKGDGWTRARRQNGEEGYVPTSYI<br>DVTLEKNSKGS                                                                                                                                                                                                                                                                                                                                                                                              |
| 3827          | A      | 2                                                                                                     | 1584                                                                                           | INPVSSAVNGEAHSSHETRGQNSNALPSVLLELL SQSCLIPAMSSYLRNDSVLDMARHVPLYRALLEL LRAIASCAAMVPLLLPLSTENGEEEEEQSECQTS VGTLLAKMKTCVDTYTNRLRSKRENVKTGVKP DASDQEPEGLTLLVPDIQKTAEIVYAATTSLRQA NQEKKLGEYSKKAAMKPKPLSVLKSLEEKYVAV MKKLQFDTFEMVSEDEDGKLGFKVNYHYMSQV KNANDANSAARARRLAQEAVTLSTSLPLSSSSSV FVRCDEERLDIMKVLITGPADTPYANGCFEFDVY FPQDYPSSPPLVNLETTGGHSVRFNPNLYNDGKV CLSILNTWHGRPEEKWNPQTSSFLQVLVSVQSLI LVAEPYFNEPGYERSRGTPSGTQSSREYDGNIRQ ATVKWAMLEQIRNPSPCFKEVIHKHFYLKRVEIM AQCEEWIADIQQYSSDKRVGRTMSHHAAALKRH TAQLREELLKLPCPEGLDPDTDDAPEVCRATTGA EETLMHDQVKPSSSKELPSDFQL |
| 3828          | A      | 1415                                                                                                  | 845                                                                                            | PRVPATLVSLDPWHCFPTAGRLAGSTWVPPACT LQLGPSSEHELDNHRAPLLSLPSQESLSFTPWYLV ACKPLFHIFCPLFACFMQEGKVQYLFLHLSHMRL LNYYFFPFLAPESLMQALEDLDYLAALDNDGNL SEFGIIMSEFPLDPQLSKSILASCEFDCVDEVLTIA AMVTGILNDYSFSFFANLH                                                                                                                                                                                                                                                                                                                                                            |
| 3829          | A      | 199                                                                                                   | 683                                                                                            | VDHTPVLSKPQCFSSVKWGATLSARSQKTSGIGR<br>LMVHVIEATELKACKPNGKSNPYCEISMGSQSYT<br>TRTIQDTLNPKWNFNCQFFIKDLYQDVLCLTLFD<br>RDQFSPDDFLGRTEIPVAKIRTEQESKGPMTRRLL<br>LHEVPTGEVWVRFDLQLFEQKTLL                                                                                                                                                                                                                                                                                                                                                                              |
| 3830          | A      | 1747                                                                                                  | 404                                                                                            | RKMMEESGIETTPPGTPPNPAGLAATAMSSTPV PLAATSSFSSPNVSSMESFPPLAYSTPQPPLPPVRP SAPLPFVPPPAVPSVPPLVTSMPPPVSPSTAAAFG NPPVSHFPPSTSAPNTLLPAPPSGPPISGFSVGSTY DITRGHAGRAPQTPLMPSFSAPSGTGLLPTPITQQ ASLTSLAQGTGTTSAITFPEEQEDPRITRGQDEAS AGGIWGFIKGVAGNPMVKSVLDKTKHSVESMIT TLDPGMAPYIKSGGELDIVVTSNKEVKVAAVRD AFQEVFGLAVVVGEAGQSNIAPQPVGYAAGLKG AQERIDSLRRTGVIHEKQTAVSVENFIAELLPDK WFDIGCLVVEDPVHGIHLETFTQATPVPLEFVQQ AQSLTPQDYNLRWSGLLVTVGEVLEKSLLNVSR TDWHMAFTGMSRRQMIYSAARAIAGMYKQRLP PRTV                                                                                    |
| 3831          | A      | 5                                                                                                     | 674                                                                                            | FWTRSAWHEGLQQMKANDPSLQEVNLYNIKNIP IPTLREFAKALETNTHVKKFSLAATRSNDPVAIAF ADMLKVNTTLTSLNIESHFITGTGILALVEALKEN DTLTEIKIDNQRQQLGTAVEMEIAQMLEENSRIL KFGYQFTKQGPRTRVAAAITKNNDLAWQKDTQ EQTSIWQVVSQSIAGFNPQFEVQGQNARSWMEE LGKAFHQFVRRELKQTEGKLP                                                                                                                                                                                                                                                                                                                          |
| 3832          | A      | 164                                                                                                   | 782                                                                                            | EPWVPMDVAESPERDPHSPEDEEQPQGLSDDDIL<br>RDSGSDQDLDGAGVRASDLEDEESAARGPSQEE                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        |

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|               |        |                                                                                                       |                                                                                                | EDNHSDEEDRASEPKSQDQDSEVNELSRGPTSSP<br>CEEEGDEGEEDRTSDLRDEASSVTRELDEHELDY<br>DEEVPEEPAPAVQEDEAEKAGAEDDEEKGEGTP<br>REEGKAGVQSVGEKESLEAAKEKKKEDDDGEID<br>DEEMY                                                                                                                                                                                                                                                                                                                                                                                          |
| 3833          | A      | 122                                                                                                   | 1676                                                                                           | SQPPHFTQKMNENKDTDSKKSEEYEDDFEKDLE WLINENEKSDASIIEMACEKEENINQDLKENETV MEHTKRHSDPDKSLQDEVSPRRNDIISVPGIQPLD PISDSDSENSFQESKLESQKDLEEEDEEVRRYIM EKIVQANKLLQNQEPVNDKRERKLKFKDQLVDL EVPPLEDTTTSKNYFENERNMFGKLSQLCISNDF GQEDVLLSLTNGSCEENKDRTILVERDGKFELLN LQDIASQGFLPPINNANSTENDPQQLLPRSSNSSV SGTKKEDSTAKIHAVTHSSTGEPLAYIAQPPLNR KTCPSSAVNSDRSKGNGKSNHRTQSAHISPVTST YCLSPRQKELQKQLEEKREKLKREEERRKIEEEK EKKRENDIVFKAWLQKKREQVLEMRRIQRAKEI EDMNSRQENRDPQQAFRLWLKKKHEEQMKERQ TEELRKQEECLFFLKGTEGRERAFKQWLRRKRM EKMAEQQAVRERTRQLRLEAKRSKQLQHHLYM SEAKPFRFTDHYN |
| 3834          | A      | 575                                                                                                   | 774 ·                                                                                          | RSRTEELSNSGILKAMSKDLVTFGDVAVNFSQEE<br>WEWLNPAQRNLYRKVMLENYRSLVSLGKDMSP                                                                                                                                                                                                                                                                                                                                                                                                                                                                               |
| 3835          | A      | 2                                                                                                     | 100                                                                                            | ASDFYLRYYVGHKGKFGHEFLEFEFRPDGVYV                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     |
| 3836          | A .    | 91                                                                                                    | 749                                                                                            | RPTPGHGDFWMQPLTKDAGMSLSSVTLASALQV<br>RGEALSEEIWSLLFLAAEQLLEDLRNDSSDYVV<br>CPWSALLSAAGSLSFQGRVSHIEAAPFKAPELLQ<br>GQSEDEQPDASQMHVYSLGMTLYWSAGFHVPP<br>HQPLQLCEPLHSILLTMCEDQPHRRCTLQSVLEA<br>CRVHEKEVSVYPAPAGLHIRRLVGLVLGTISEVS<br>REPCFSSSSCWSCVAIKI                                                                                                                                                                                                                                                                                                   |
| 3837          | A      | 3                                                                                                     | 1214                                                                                           | SLGCTNSARGKGQDDEVRTLMANGAPFTTDWFS KLRVSCGYIGDNCKNGADVNAKDMLKMTALH WATERHHRDVVELLIKYGADVHAFSKFDKSAFD IALEKNNAEILVILQEAMQNQVNVNPERANPVTD PVSMAAPFIFTSGEVVNLASLISSTNTKTTSGDPH ASTVQFSNSTTSVLATLAALAEASVPLSNSHRAT ANTEEIIEGNSVDSSIQQVMGSGGQRVITIVTDGV PLGNIQTSIPTGGIGHPFIVTVQDGQQVLTVPAGK VAEETVIKEEEEEKLPLTKKPRIGEKTNSVEESKE GNERELLQQQLQEANRRAQEYRHQLLKKEQEAE QYRLKLEAIARQQPNGVDFTMVEEVAEVDAVV VTEGELEERETKVTGSAGATGPPTRVSMATVSS                                                                                                                       |
| 3838          | A      | 1                                                                                                     | 1332                                                                                           | MIEDNKENKDHSLERGRASLIFSLKNEVGGLIKA LKIFQEKHVNLLHIESRKSKRRNSEFEIFVDCDIN REQLNDIFHLLKSHTNVLSVNLPDNFTLKEDGME TVPWFPKKISDLDHCANRVLMYGSELDADHPGF KDNVYRKRRKYFADLAMNYKHGDPIPKVEFTEE EIKTWGTVFQELNKLYPTHACREYLKNLPLLSKY CGYREDNIPQLEDVSNFLKERTGFSIRPVAGYLSP RDFLSGLAFRVFHCTQYVRHSSDPFYTPEPDTCH ELLGHVPLLAEPSFAQFSQEIGLASLGASEEAVQ KLATCYFFTVEFGLCKQDGQLRVFGAGLLSSISE LKHALSGHAKVKPFDPKITCKQECLITTFQDVYF VSESFEDAKEKMREFTKTIKRPFGVKYNPYTRSI                                                                                                                  |

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|               |        |                                                                                                       |                                                                                                | QILKDTKSITSAMNELQHDLDVVSDALAKVSRKP<br>SI                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            |
| 3839          | A      | 3093                                                                                                  | 520                                                                                            | MVNFTVDQIRAIMDKKANIRNMSVIAHVDHGKS TLTDSLVCKAGIIASARAGETRFTDTRKDEQERCI TIKSTAISLFYELSENDLNFIKQSKDGAGFLINLID SPGHVDFSSEVTAALRVTDGALVVVDCVSGVCV QTETVLRQAIAERIKPVLMMNKMDRALLELQLE PEELYQTFQRIVENVNVIISTYGEGESGPMGNIMI DPVLGTVGFGSGLHGWAFTLKQFAEMYVAKFA AKGEGQLGPAERAKKVEDMMKKLWGDRYFDP ANGKFSKSATSPEGKKLPRTFCQLILDPIFKVFDA IMNFKKEETAKLIEKLDIKLDSEDKDKEGKPLLK AVMRRWLPAGDALLQMITIHLPSPVTAQKYRCE LLYEGPPDDEAAMGIKSCDPKGPLMMYISKMVP TSDKGRFYAFGRVFSGLVSTGLKVRIMGPNYTPG KKEDLYLKPIQRTILMMGRYVEPIEDVPCGNIVG LVGVDQFLVKTGTITTFEHAHNMRVMKFSVSPV VRVAVEAKNPADLPKLVEGLKRLAKSDPMVQCI IEESGEHIIAGAGELHLEICLKDLEEDHACIPIKKS DPVVSYRETVSEESNVLCLSKSPNKHNRLYMKA RPFPDGLAEDIDKGEVSARQELKQRARYLAEKY EWDVAEARKIWCFGPDGTGPNILTDITKGVQYL NEIKDSVVAGFQWATKEGALCEENMRGVRFDV HDVTLHADAIHRGGGQIIPTARRCLYASVLTAQP RLMEPIYLVEIQCPEQVVGGIYGVLNRKRGHVFE ESQVAGTPMFVVKAYLPVNESFGFTADLRSNTG GQAFPQCVFDHWQILPGDPFDNSSRPSQVVAETR KRKGLKEGIPALDNFLDKL |
| 3840          | A      | 2                                                                                                     | 753                                                                                            | SSTRSRDFCCSEAIQGSLTRRERRASGVRTRRSQG<br>SSAMASKILLNVQEEVTCPICLELLTEPLSLDCGH<br>SLCRACITVSNKEAVTSMGGKSSCPVCGISYSFE<br>HLQANQHLANIVERLKEVKLSPDNGKKRDLCDH<br>HGEKLLLFCKEDRKVICWLCERSQEHRGHHTVL<br>TEEVFKECQEKLQAVLKRLKKEEEEAEKLEADIR<br>EEKTSWKYQVQTERQRIQTEFDQLRSILNNEEQR<br>ELQRLEEEEKKT                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              |
| 3841          | A      | 2                                                                                                     | 405                                                                                            | GKAFSCFTYLSQHRRTHMAEKPYECKTCKKAFS<br>HFGNLKVHERIHTGEKPYECKECRKAFSWLTCL<br>LRHERIHTGKKSYECQQCGKAFTRSRFLRGHEKT<br>HTGEKMHECKECGKALSSLSSLHRHKRTHWRDT<br>L                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              |
| 3842          | A      | 311                                                                                                   | 88                                                                                             | AVLKNMAPMTALGLLDLHILNLILFLSAGEDFTS<br>VVSEIMMYILLVFLTLWLLIEMIYCYRKVSKAEE<br>AAQENA                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  |
| 3843          | A      | 3                                                                                                     | 1175                                                                                           | APIRNSRIDDFVRRVESKATSARCGLWGSGPRRR PASGMFRGLSSWLGLQQPVAGGGQPNGDAPPEQ PSETVAESAEEELQQAGDQELLHQAKDFGNYLF NFASAATKKITESVAETAQTIKKSVEEGKIDGIID KTIIGDFQKEQKKFVEEQHTKKSEAAVPPWVDT NDEETIQQQILALSADKRNFLRDPPAGVQFNFDF DQMYPVALVMLQEDELLSKMRFALVPKLVKEE VFWRNYFYRVSLIKQSAQLTALAAQQQAAGKEE KSNGREQDLPLAEAVRPKTPPVVIKSQLKTQEDE EEISTSPGVSEFVSDAFDACNLNQEDLRKEMEQL VLDKKQEETAVLEEDSADWEKELQQELQEYEV                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           |

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|               | <u> </u> | sequence                                                                                     |                                                                                                | TAMPODAND DE MADALERA AL OPENA                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      |
| L             |          |                                                                                              |                                                                                                | VTESEKRDENWDKEIEKMLQEEN                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             |
| 3844          | A ·      | 798                                                                                          | 148                                                                                            | LPPAQIPEAWLLLANVVVVLILVPLKDRLIDPLLL<br>RCKLLPSALQKMALGMFFGFTSVIVAGVLEMER<br>LHYIHHNETVSQQIGEVLYNAAPLSIWWQIPQYL<br>LIGISEIFASIPGLEFAYSEAPRSMQGAIMGIFFCLS<br>GVGSLLGSSLVALLSLPGGWLHCPKDFGNINNCR<br>MDLYFFLLAGIQAVTALLFVWIAGRYERASQGP                                                                                                                                                                                                                                                                                                                                                                                                                  |
|               | }        | ł                                                                                            |                                                                                                | ASHSRFSRDRG                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         |
| 3845          | A        | 3                                                                                            | 1934                                                                                           | PEDSAPQYSRLFPNASQHITPSYNYAPNPDKHWI MRYTGPMKPIHMEFTNMLQRKRLQTLMSVDDS METIYNMLVETGELDNTYIVYTADHGYHIGQFG LVKGKSMPYEFDIRVPFYVRGPNVEAGCLNPHIV LNIDLAPTILDIAGLDIPADMDGKSILKLLDTERP VNRFHLKKKMRVWRDSFLVERGKLLHKRDNDK VDAQEENFLPKYQRVKDLCQRAEYQTACEQLG QKWQCVEDATGKLKLHKCKGPMRLGGSRALSN LVPKYYGQGSEACTCDSGDYKLSLAGRRKKLFK KKYKASYVRSRSIRSVAIEVDGRVYHVGLGDAA QPRNLTKRHWPGAPEDQDDKDGGDFSGTGGLP DYSAANPIKVTHRCYILENDTVQCDLDLYKSLQ AWKDHKLHIDHEIETLQNKIKNLREVRGHLKKK RPEECDCHKISYHTQHKGRLKHRGSSLHPFRKGL                                                                                                                                                         |
| 3846          | A        | 3                                                                                            | 1934                                                                                           | QEKDKVWLLREQKRKKKLRKLLKRLQNNDTCS MPGLTCFTHDNQHWQTAPFWTLGPFCACTSAN NNTYWCMRTINETHNFLFCEFATGFLEYFDLNT DPYQLMNAVNTLDRDVLNQLHVQLMELRSCKG YKQCNPRTRNMDLGLKDGGSYEQYRQFQRRKW PEMKRPSSKSLGQLWEGWEG PEDSAPQYSRLFPNASQHITPSYNYAPNPDKHWI                                                                                                                                                                                                                                                                                                                                                                                                                       |
| -             |          | ·                                                                                            |                                                                                                | MRYTGPMKPIHMEFTNMLQRKRLQTLMSVDDS METIYNMLVETGELDNTYIVYTADHGYHIGQFG LVKGKSMPYEFDIRVPFYVRGPNVEAGCLNPHIV LNIDLAPTILDIAGLDIPADMDGKSILKLLDTERP VNRFHLKKKMRVWRDSFLVERGKLLHKRDNDK VDAQEENFLPKYQRVKDLCQRAEYQTACEQLG QKWQCVEDATGKLKLHKCKGPMRLGGSRALSN LVPKYYGQGSEACTCDSGDYKLSLAGRRKKLFK KKYKASYVRSRSIRSVAIEVDGRVYHVGLGDAA QPRNLTKRHWPGAPEDQDDKDGGDFSGTGGLP DYSAANPIKVTHRCYILENDTVQCDLDLYKSLQ AWKDHKLHIDHEIETLQNKIKNLREVRGHLKKK RPEECDCHKISYHTQHKGRLKHRGSSLHPFRKGL QEKDKVWLLREQKRKKKLRKLLKRLQNNDTCS MPGLTCFTHDNQHWQTAPFWTLGPFCACTSAN NNTYWCMRTINETHNFLFCEFATGFLEYFDLNT DPYQLMNAVNTLDRDVLNQLHVQLMELRSCKG YKQCNPRTRNMDLGLKDGGSYEQYRQFQRRKW PEMKRPSSKSLGQLWEGWEG |
| 3847          | A        | 1                                                                                            | 1257                                                                                           | MVFSAVLTAFHTGTSNTTFVVYENTYMNITLPPP<br>FQHPDLSPLLRYSFETMAPTGLSSLTVNSTAVPTT<br>PAAFKSLNLPLQITLSAIMIFILFVSFLGNLVVCLM<br>VYQKAAMRSAINILLASLAFADMLLAVLNMPFA<br>LVTILTTRWIFGKFFCRVSAMFFWLFVIEGVAILL<br>IISIDRFLIIVQRQDKLNPYRAKVLIAVSWATSFCV<br>AFPLAVGNPDLQIPSRAPQCVFGYTTNPGYQAYV                                                                                                                                                                                                                                                                                                                                                                         |

| 070 -         | 1 30 1 | I D. D. C.                                                                                            | Deciles 1                                                                                      | Literature (A-Alexia C Continue Date of the                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       |
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|               |        |                                                                                                       |                                                                                                | ILISLISFIPFLVILYSFMGILNTLRHNALRIHSYPE<br>GICLSQASKLGLMGLQRPFQMSIDMGFKTRAFTT<br>ILILFAVFIVCWAPFTTYSLVATFSKHFYYQHNFF<br>EISTWLLWLCYLKSALNPLIYYWRIKKFHDACLD<br>MMPKSFKFLPQLPGHTKRRIRPSAVYVCGEHRT                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     |
|               |        |                                                                                                       |                                                                                                | VV                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                |
| 3848          | A      | 3                                                                                                     | 2827                                                                                           | SSAVAARRRSWASLVLAFLGVCLGITLAVDRS NFKTCEESSFCKRQRSIRPGLSPYRALLDSLQLGP DSLTVHLIHEVTKVLLVLELQGLQKNMTRFRIDE LEPRRPRYRVPDVLVADPPIARLSVSGRDENSVE LTMAEGPYKIILTARPFRLDLLEDRSLLLSVNARG LLEFEHQRAPRVSQGSKDPAEGDGAQPEETPRD GDKPEETQGKAEKDEPGAWEETFKTHSDSKPYG PMSVGLDFSLPGMEHVYGIPEHADNLRLKVTEG GEPYRLYNLDVFQYELYNPMALYGSVPVLLAHN PHRDLGIFWLNAAETWVDISSNTAGKTLFGKMM DYLQGSGETPQTDVRWMSETGIIDVFLLLGPSISD VFRQYASLTGTQALPPLFSLGYHQSRWNYRDEA DVLEVDQGFDDHNLPCDVIWLDIEHADGKRYFT WDPSRFPQPRTMLERLASKRRKLVAIVDPHIKVD SGYRVHEELRNLGLYVKTRDGSDYEGWCWPGS AGYPDFTNPTMRAWWANMFSYDNYEGSAPNLF VWNDMNEPSVFNGPEVTMLKDAQHYGGWEHR DVHNIYGLYVHMATADGLRQRSGGMERPFVLA RAFFAGSQRFGAVWTGDNTAEWDHLKISIPMCL SLGLVGLSFCGADVGGFFKNPEPELLVRWYQMG AYQPFFRAHAHLDTGRREPWLLPSQHNDIIRDAL GQRYSLLPFWYTLLYQAHREGIPVMRPLWVQYP QDVTTFNIDDQYLLGDALLVHPVSDSGAHGVQV YLPGQGEVWYDIQSYQKHHGPQTLYLPVTLSSIP VFQRGGTIVPRWMRVRRSSECMKDDPITLFVALS PQGTAQGELFLDDGHTFNYQTRQEFLLRRFSFSG NTLVSSSADPEGHFETPIWIERVVIIGAGKPAAVV LQTKGSPESRLSFQHDPETSVLVLRKPGINVASD |
| 3849          | A      | 1                                                                                                     | 1717                                                                                           | WSIHLR  RARNARGCWGVCRSGFSSAVCGAARMEQVAEG ARVTAVPVSAADSTEELAEVEEGVGVVGEDNDA AARGAEAFGDSEEDGEDVFEVEKILDMKTEGGK VLYKVRWKGYTSDDDTWEPEIHLEDCKEVLLEF RKKIAENKAKAVRKDIQRLSLNNDIFEANSDSDQ QSETKEDTSPKKKKKKLRQREEKSPDDLKKKKA KAGKLKDKSKPDLESSLESLVFDLRTKKRISEAK EELKESKKPKKDEVKETKELKKVKKGEIRDLKT KTREDPKENRKTKKEKFVESQVESESSVLNDSPF PEDDSEGLHSDSREEKQNTKSARERAGQDMGLE HGFEKPLDSAMSAEEDTDVRGRRKKKTPRKAED TRENRKLENKNAFLEKKTVPKKQRNQDRSKSAA ELEKLMPVSAQTPKGRRLSGEERGLWSTDSAEE DKETKRNESKKPKKDEVKETKELKKVKKGEIRD LKTKTREDPKENRKTKKEKFVESQVESESSVLND SPFPEDDSEGLHSDSREEKQNTKSARERAGQDM GLEHGFEKPLDSAMSAEEDTDVRGRRKKKTPRK AEDTRENRKLENKNAFLEKKTVPKKQRNQDRSK SAAELEKLMPVSAQTPKGRRLSGEERGLWSTDS AEEDKETKRNESKKPKKDEVKETKELKKVKKGE                                                                                                                                                                                                                                                                                |

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|---------------|--------|-------------------------------------------------------------------------------------------------------|------------------------------------------------------------------------------------------------|-------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
|               |        |                                                                                                       |                                                                                                | IRDLKTKTREDPKENRKTKKEKFVESQVESESSV<br>LNDSPFPED/RQ*RATFRQQREEKSPDDLKKKKA<br>KAGKLKDKSKPDLESSLESLVFDLRTKKRISEAK<br>EELKESKKPK                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            |
| 3850          | A .    | 1113                                                                                                  | 3975                                                                                           | PAAAAAAAAAAAAAAAGRGPSFTPCFSPSLAVEPS RRTRLGSDPAQAMAGNVKKSSGAGGGSGGS GSGGLIGLMKDAFQPHHHHHHHHLSPHPPGTVDK KMVEKCWKLMDKVVRLCQNPKLALKNSPPYIL DLLPDTYQHLRTILSRYEGKMETLGENEYFRVF MENLMKKTKQTISLFKEGKERMYEENSQPRRNL TKLSLIFSHMLAELKGIFPSGLFQGDTFRITKADA AEFWRKAFGEKTIVPWKSFRQALHEVHPISSGLE AMALKSTIDLTCNDYISVFEFDIFTRLFQPWSSLL RNWNSLAVTHPGYMAFLTYDEVKARLQKFIHKP GSYIFRLSCTRLGQWAIGYVTADGNILQTIPHNKP LFQALIDGFREGFYLFPDGRNQNPDLTGLCEPTP QDHIKVTQEQYELYCEMGSTFQLCKICAENDKD VKIEPCGHLMCTSCLTSWQESEGQGCPFCRCEIK GTEPIVVDPFDPRGSGSLLRQGAEGAPSPNYDDD DDERADDTLFMMKELAGAKVERPPSPFSMAPQA SLPPVPPRLDLLPQRVCVPSSASALGTASKAASGS LHKDKPLPVPPTLRDLPPPPPPDRPYSVGAESRPQ RRPLPCTPGDCPSRDKLPPVPSSRLGDSWLPRPIP KVPVSAPSSSDPWTGRELTNRHSLPFSLPSQMEP RPDVPRLGSTFSLDTSMSMNSSPLVGPECDHPKI KPSSSANAIYSLAARPLPVPKLPPGEQCEGEEDTE YMTPSSRPLRPLDTSQSSRACDCDQQIDSCTYEA MYNIQSQAPSITESSTFGEGNLAAAHANTGPEES ENEDDGYDVPKPPVPAVLARRTLSDISNASSS/FG LFVLERDP*PQNVTEGSQVPERPPKPFPRRINSER KAGSCQQGSGPAASAATA\SPQLSSEIENLMSQG YSYQDIQKALVIAQNNIEMAKNILREFVSISSPAH VAT |
| 3851          | A      |                                                                                                       | 2781                                                                                           | GRVGSMDGAMGPRGLLLCMYLVSLLILQAMPA LGSATGRSKSSEKRQAVDTAVDGVFIRSLKVNC KVTSRFAHYVVTSQVVNTANEAREVAFDLEIPK TAFISDFAVTADGNAFIGDIKDKVTAWKQYRKA AISGENAGLVRASGRTMEQFTIHLTVNPQSKVTF QLTYEEVLKRNHMQYEIVIKVKPKQLVHHFEIDV DIFEPQGISKLDAQASFLPKELAAQTIKKSFSGKK GHVLFRPTVSQQQSCPTCSTSLLNGHFKVTYDVS RDKICDLLVANNHFAHFFAPQNLTNMNKNVVFV IDISGSMRGQKVKQTKEALLKILGDMQPGDYFD LVLFGTRVQSWKGSLVQASEANLQAAQDFVRGF SLDEATNLNGGLLRGIEILNQVQESLPELSNHASI LIMLTDGDPTEGVTDRSQILKNVRNAIRGRFPLY NLGFGHNVDFNFLEVMSMENNGRAQRIYEDHD ATQQLQGFYSQVAKPLLVDVDLQYPQDAVLALT QNHHKQYYEGSEIVVAGRIADNKQSSFKADVQA HGEGQEFSITCLVDEEEMKKLLRERGHMLENHV ERLWAYLTIQELLAKRMKVDREVRANLSSQALR MSLDYGFVTPLTSMSIRGMADQDGLKPTIDKPSE DSPPLEMLGPRRTFVLSALQPSPTHSSSNTQRLPD RVTGVDTDPHFIIHVPQKEDTLCFNINEEPGVILS LVQDPNTGFSVNGQLIGNKARSPGQHDGTYFGR                                                                                                                                                                                                                                  |

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|               | No. 22 - 2 | Dung!-4-3                                                                                             | Dunglasad                                                                                      | Amino said seguence (AmAlanine C-Custoine D-A-said Asid                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           |
|---------------|------------|-------------------------------------------------------------------------------------------------------|------------------------------------------------------------------------------------------------|-----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
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|               |            |                                                                                                       |                                                                                                | S\GPGAPTPAAPPQLARMAWAFSLSAASTPAVSP<br>STSPSAVEGSPATILPLASSPPPRTTP*LPLSELTV*<br>RPQELVRGRGCLGPGAPTPAAPPQLARMAWAFS<br>LSAASTPAVSPSTSPSAVEGSPATILPLASSPPPRT<br>TP                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    |
| 3855          | A          | 1                                                                                                     | 772                                                                                            | FRGGDGAPGVLKPGNPLPFPLPPLQYPPPSTLSHS DNLAMTSRSTARPNGQPQASKICQFKLVLLGESA VGKSSLVLRFVKGQFHEYQESTIGAAFLTQSVCL DDTTVKFEIWDTAGQERYHSLAPMYYRGAQAAI VVYDITNQETFARAKTWVKELQRQASP\SIVVGL AGNKADLANKRMVEYEEAQAYADDNSLLFMET SAKTAMNVNDLFL\AIA*EVAKRVNPQNLG\G\A AGRSRGVDLHEQS\QQNKSQCCSN                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       |
| 3856          | A          | 2815                                                                                                  | 352                                                                                            | LGLEAAARPRPGGPAAMQDGNFLLSALQPEAGV CSLALPSDLQLDRRGAEGPEAERLRAARVQEQV RARLLQLGQQPRHNGAAEPEPEAETARGTSRGQ YHTLQAGFSSRSQGLSGDKTSGFRPIAKPAYSPA SWSSRSAVDLSCSRRLSSAHNGGSAFGAAGYGG AQPTPPMPTRPVSFHERGGVGSRADYDTLSLRSL RLGPGGLDDRYSLVSEQLEPAATSTYRAFAYER QASSSSRAGGLDWPEATEVSPSRTIRAPAVRTL QRFQSSHRSRGVGGAVPGAVLEPVARAPSVRSLS LSLADSGHLPDVHGFNSYGSHRTLQRLSSGFDDI DLPSAVKYLMASDPNLQVLGAAYIQHKCYSDAA AKKQARSLQAVPRLVKLFNHANQEVQRHATGA MRNLIYDNADNKLALVEENGIFELLRTLREQDDE LRKNVTGILWNLSSSDHLKDRLAKKTPLE\QLT\D LGV*APLSGAGGPP\LIQQNASEAEIFYNATGFPR NLSSASQATRQKMRECHGLVDALVTSINHALDA GKCEDKSVENAVCVLRNLSYRLYDEMPPSALQR LEGRGRRDLAGAPPGEVVGCFTPQSRRLRELPLA ADALTFAEVSKDPKGLEWLWSPQIVGLYNRLLQ RCELNRHTTEAAAGALQNITGG\DPRGPGGLSRL ALEQERILNPLLDRVRTADHHQLRSLTGLIRNLS RNARNKDEMSTKVV\SHLI\EKLPGSVGEKSPPAE VLV\NI\IAVFNNLGWLASPI/ALARDLLYFDGLRK LIFIKKKRDSPDSEKSSRAASSLLANLWQYNKLH RDFRAKGYRKEDFLGP |
| 3857          | A .        | 1034                                                                                                  | 204                                                                                            | VAVTLLSQLPSAIQRTAAWEMRAPLTFRVPLALD LIKPEHCTVNVDNSLSIPVIAAELVVRKPSEKGM QQKKKTKDLGFRAGKESKTEWRK*GLQDMASQ MFALPLK*PVTAAFHDSSMPSSLLQIEMEQLFLE ARLQ/PDSKSEARRNQCDSMLLRNQQLCSTCQE MKMVQPRTMKIPDDPKASFENCMSYRMSLHQP KFQTTPEPFHDDIPTENIHLQNL/PILGPRTAVFHG LLTEAYKTLKERQRSSLPRKEPIGKTTEAVSGRSS SPPRLPERK                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   |
| 3858          | A          |                                                                                                       | 3469                                                                                           | SHQEIEQNSAMAPRKRGGRGISFIFCCFRNNDHPE ITYRLRNDSNFALQTMEPALPMPPVEELDVMFSE LVDELDLTDKHREAMFALPAEKKWQIYCSKKK DQEENKGATSWPEFYIDQLNSMAARKSLLALEK EEEEERSKTIESLKTALRTKPMRFVTRFIDLDGLS CILNFLKTMDYETSESRIHTSLIGCIKALMNNSQG RAHVLAHSESINVIAQSLSTENIKTKVAVLEILGA VCLVPGGHKKVLQAMLHYQKYASERTRFQTLIN                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           |

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|---------------|--------|-------------------------------------------------------------------------------------------------------|------------------------------------------------------------------------------------------------|--------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
|               |        |                                                                                                       |                                                                                                | DLDKSTGRYRDEVSLKTAIMSFINAVLSQGAGVE SLDFRLHLRYEVFLMLGIHPVMDKLRKHENSTLD RHLDFFEMLRNEDELEFAKRFELVHIDTKSATQM FELTRKRLTHSEAYPHFMSILHHCLQMPYKRSGN TVQYWLLLDRIIQQIVIQNDKGQDPDSTPLENFNI KNVVRMLVNENEVKQWKEQAEKMRKEHNELQ QKLEKKERECDAKTQEKEEMMQTLNKMKEKLE KETTEHKQVKQQVADLTAQLHELSRRAVCASIP GGPSPGAPGGPFPSSVPGSLLPPPPPPPPLPGGMLPP PPPPLPPGGPPPPGPPLGAIMPPPGAPMGLALK KKSIPQPTNALKSFNWSKLPENKLEGTVWTEIDD TKVFKILDLEDLERTFSAYQRQQDFFVNSNSKQK EADAIDDTLSSKLKVKELSVIDGRRAQNCNILLS RLKLSNDEIKRAILTMDEQEDLPKDMLEQLLKFV PEKSDIDLLEEHKHELDRMAKADRFLFEMSRINH YQQRLQSLYFKKKFAERVAEVKPKVEAIRSGSEE VFRSGALKQLLEVVLAFGNYMNKGQRGNAYGF KISSLNKIADTKSSIDKNITLLHYLITIVENKYPSV LNLNEELRDIPQAAKVNMTELDKEISTLRSGLKA VETELEYQKSQPPQPGDKFVSVVSQFITVASFSFS DVEDLLAEAKDLFTKAVKHFGEEAGKIQPDEFF GIFDQFLQAVSEAKQENENMRKKKEEEERRARM EAQLKEQRERERKMRKAKENSEESGEFDDLVSA LRSGEVFDKDLSKLKRNRKRITNQMTDSSRERPI TKLNF |
| 3859          | A      | 1279                                                                                                  | 141                                                                                            | RVEHLSEFLVDIKPSLTFDVIPLLDPYGPAGSDPS LEFLVVSEETYRGGMAINRFRLENDLEELALYQI QLLKDLRHTENEEDKVSSSSFRQRMLGNLLRPPY ERPELPTCLYVIGLTGISGSGKSSIAQRLKGLGAF VIDSDHLGHRAYAPGGPAYQPVVEAFGTDILHK DGIINRKVLGSRVFGNKKQLKILTDIMWPIIAKLA REEMDRAVAEGKRVCVIDAAVLLEAGWQNLVH EVWTAVIPETEAVRRIVERDGLSEAAAQSRLQSQ MSGQQLVEQSHVVLST\CGSRISPNARWRKPGPS CRSAFPRLIRPSTEKFSVGPDWLLELTSDPVVRN GGLDAHPGSGPEVQAILCRTWPGLVDTGSLPNTL VFGQH                                                                                                                                                                                                                                                                                                                                                                                                                                                                   |
| 3860          | A      | 1                                                                                                     | 3881                                                                                           | MGQKSVGASYVQIPLVPPLSRHPKGLGHEDRWS SYCLSSLAAQNICTSKLHCPAAPEHTDPSEPRGSV SCCSLLRGLSSGWSSPLLPAPVCNPNKAIFTVDA KTTEILVANDKACGLLGYSSQDLIGQKLTQFFLR SDSDVVEALSEEHMEADGHAAVVFGTVVDIISRS GEKIPVSVWMKRMRQERRLCCVVVLEPVERVST WVAFQSDGTVTSCDSLFAHLHGYVSGEDVAGQ HITDLIPSVQLPPSGQHIPKNLKIQRSVGRARDGT TFPLSLKLKSQPSSEEATTGEAAPVSGYRASVWV FCTISGLITLLPDGTIHGINHSFALTLFGYGKTELL GKNITFLIPGFYSYMDLAYNSSLQLPDLASCLDV GNESGCGERTLDPWQGQDPAEGGQDPRINVVLA GGHVVPRDEIRKLMESQDIFTGTQTELIAGGQLL SCLSPQPAPGVDNVPEGSLPVHGEQALPKDQQIT ALGREEPVAIESPGQDLLGESRSEPVDVKPFASCE DSEAPVPAEDGGSDAGMCGLCQKAQLERMGVS GPSGSDLWAGAAVAKPQAKGQLAGGSLLMHCP CYGSEWGLWWRSQDLAPSPSGMAGLSFGTPTLD                                                                                                                                                                                                                         |

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Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, Predicted end SEQ ID Method Predicted nucleotide E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, beginning NO: I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, nucleotide location N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, corresponding location T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, corresponding to last amino X=Unknown, \*=Stop codon, /=possible nucleotide deletion, acid residue of to first amino acid residue of peptide \=possible nucleotide insertion peptide sequence sequence **EPWLGVENDREELQTCLIKEQLSQLSLAGALDVP** HAELVPTECQAVTAPVSSCDLGGRDLCGGCTGS SSACYALATDLPGGLEAVEAQEVDVNSFSWNLK ELFFSDOTDOTSSNCSCATSELRETPSSLAVGSDP DVGSLOEOGSCVLDDRELLLLTGTCVDLGQGRR FRESCYGHDPTEPLEVCLYSSEHYAASDRESPGH VPSTLDAGPEDTCPSAEEPRLNVQVTSTPVIVMR GAAGLQREIQEGAYSGSCYHRDGLRLSIQFEVRR VELOGPTPLFCCWLVKDLLHSQRDSAARTRLFL **ASLPGSTHSTAAELTGPSLVEVLRARPWFEEPPK** AVELEGLAACEGEYSQKYSTMSPLGSGAFGFVW TAVDKEKNKEVVVKFIKKEKVLEDCWIEDPKLG KYTLEIAILSRVEHANIIKVLDIFENQGFFQLVME KHGSGLDLFAFIDRHPRLDEPLASYIFRQVRAG\Q SRLVSAVGYLRLKDIIHRDIKDENIVIAEDFTIKLI DFGSAAYLERGKLFYTFCGTIEYCAPEVLMGNPY RGPELEMWSLGVTLYTLVFEENPFCELEETVEAA IHPPYLVSKELMSLVSGLLQPVPERRTTLEKLVT **DPWVTQPVNLADYTWEEVFRVNKPESGVLSAAS** LEMGNRSLSDVAQAQELCGGPVPGEAPNGQGCL **HPGDPRLLTS** MGQKSVGASYVQIPLVPPLSRHPKGLGHEDRWS 3861 3881 SYCLSSLAAQNICTSKLHCPAAPEHTDPSEPRGSV SCCSLLRGLSSGWSSPLLPAPVCNPNKAIFTVDA KTTEILVANDKACGLLGYSSQDLIGQKLTQFFLR SDSDVVEALSEEHMEADGHAAVVFGTVVDIISRS GEKIPVSVWMKRMRQERRLCCVVVLEPVERVST WVAFQSDGTVTSCDSLFAHLHGYVSGEDVAGQ HITDLIPSVQLPPSGQHIPKNLKIQRSVGRARDGT TFPLSLKLKSQPSSEEATTGEAAPVSGYRASVWV FCTISGLITLLPDGTIHGINHSFALTLFGYGKTELL GKNITFLIPGFYSYMDLAYNSSLQLPDLASCLDV GNESGCGERTLDPWQGQDPAEGGQDPRINVVLA GGHVVPRDEIRKLMESQDIFTGTQTELIAGGQLL SCLSPQPAPGVDNVPEGSLPVHGEQALPKDQQIT ALGREEPVAIESPGQDLLGESRSEPVDVKPFASCE DSEAPVPAEDGGSDAGMCGLCQKAQLERMGVS GPSGSDLWAGAAVAKPQAKGQLAGGSLLMHCP CYGSEWGLWWRSQDLAPSPSGMAGLSFGTPTLD **EPWLGVENDREELQTCLIKEQLSQLSLAGALDVP HAELVPTECQAVTAPVSSCDLGGRDLCGGCTGS** SSACYALATDLPGGLEAVEAQEVDVNSFSWNLK ELFFSDQTDQTSSNCSCATSELRETPSSLAVGSDP DVGSLQEQGSCVLDDRELLLLTGTCVDLGQGRR FRESCVGHDPTEPLEVCLVSSEHYAASDRESPGH **VPSTLDAGPEDTCPSAEEPRLNVQVTSTPVIVMR** GAAGLOREIQEGAYSGSCYHRDGLRLSIQFEVRR VELQGPTPLFCCWLVKDLLHSQRDSAARTRLFL ASLPGSTHSTAAELTGPSLVEVLRARPWFEEPPK **AVELEGLAACEGEYSQKYSTMSPLGSGAFGFVW** TAVDKEKNKEVVVKFIKKEKVLEDCWIEDPKLG KVTLEIAILSRVEHANIIKVLDIFENQGFFQLVME KHGSGLDLFAFIDRHPRLDEPLASYIFRQVRAG\Q SRLVSAVGYLRLKDIIHRDIKDENIVIAEDFTIKLI DFGSAAYLERGKLFYTFCGTIEYCAPEVLMGNPY

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|---------------|--------|-------------------------------------------------------------------------------------------------------|------------------------------------------------------------------------------------------------|-----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
|               |        |                                                                                                       |                                                                                                | RGPELEMWSLGVTLYTLVFEENPFCELEETVEAA IHPPYLVSKELMSLVSGLLQPVPERRTTLEKLVT DPWVTQPVNLADYTWEEVFRVNKPESGVLSAAS LEMGNRSLSDVAQAQELCGGPVPGEAPNGQGCL HPGDPRLLTS                                                                                                                                                                                                                                                                                                                                                                                                                                          |
| 3862          | A      | 399                                                                                                   | 2069                                                                                           | TMDRSKRNSIAGFPPRVE\RLEEFEGGGGGEGNV SQVGRVWPSSYRALISAFFRLTRLDDFTCEKIGSG FFSEVFKVRHRASGQVMALKMNTLSSNRANML KEVQLMNRLSHPNILRYINSGNLEQLLDSNLHLP WTVRVKLAYDIAVGLSYLHFKGIFHRDLTSKNC LIKRDENGYSAVVADFGLAEKIPDVSMGSEKLA VVGSPFWMAPEVLRDEPYNEKADVFSYGIILCEII ARIQADPDYLPRTENFGLDYDAFQHMVGDCPPD FLQLTFNCCNMDPKLRPSFVEIGKTLEEILSRLQE EEQERDRKLQPTARGLLEKAPGVKRLSSLDDKIP HKSPCPRRTIWLSRSQSDIFSRKPPRTVSVLDPYY RPRDGAARTPKVNPFSARQDLMGGKIKFFDLPSK SVISLVFDLDAPGPGTMPLADWQEPLAPPIRRWR SLPGSPEFLHQEACPFVGREESLSDGPPPRLSSLK YRVKEIPPFRASALPAAQAHEAMDCSILQEENGF GSRPQGTSPCPAGASEEMEVEERPAGSTPATFSTS GIGLQTQGKQDG |
| 3863          | A .    |                                                                                                       | 2069                                                                                           | TMDRSKRNSIAGFPPRVE/RLEEFEGGGGGEGNV SQVGRVWPSSYRALISAFFRLTRLDDFTCEKIGSG FFSEVFKVRHRASGQVMALKMNTLSSNRANML KEVQLMNRLSHPNILRYINSGNLEQLLDSNLHLP WTVRVKLAYDIAVGLSYLHFKGIFHRDLTSKNC LIKRDENGYSAVVADFGLAEKIPDVSMGSEKLA VVGSPFWMAPEVLRDEPYNEKADVFSYGIILCEII ARIQADPDYLPRTENFGLDYDAFQHMVGDCPPD FLQLTFNCCNMDPKLRPSFVEIGKTLEEILSRLQE EEQERDRKLQPTARGLLEKAPGVKRLSSLDDKIP HKSPCPRRTIWLSRSQSDIFSRKPPRTVSVLDPYY RPRDGAARTPKVNPFSARQDLMGGKIKFFDLPSK SVISLVFDLDAPGPGTMPLADWQEPLAPPIRRWR SLPGSPEFLHQEACPFVGREESLSDGPPPRLSSLK YRVKEIPPFRASALPAAQAHEAMDCSILQEENGF GSRPQGTSPCPAGASEEMEVEERPAGSTPATFSTS GIGLQTQGKQDG |
| 3864          | A      | 3                                                                                                     | 911                                                                                            | SWNMDSDSCAAAFHPEEYSPSCKRRTVEDFNK FCTFVLAYAGYIPYPKEELPLRSSPSPANSTAGTI DSDGWDAGFSDIASSVPLPVSDRCFSHLQPTLLQ RAKPSNFLLDRKKTDKLKKKKKKRRDSDAPGK EGYRGGLLKLEAADPYVETPTSPTLQDIPQAPSD PCSGWDSDTPSSGSCATVSPDQVKEIKTEGKRTI VR/QEAQLMARNDGNFSSLLESIFPS\DDDSWDLV TCFCMKPFAGRPMIECNECHTWIHLSCAKIRKSN VPEVFVCQKCRDSKFDIRRSNRSRTGSRKLFLD                                                                                                                                                                                                                                                                       |
| 3865 .        | A      | 3                                                                                                     | 3573                                                                                           | QERLRSRSRPDRAAREAGSARGRQPKRTERVEQ<br>FLTIARRRGRRSMPVSLEDSGEPTSCPATDAETAS<br>EGSVESASETRSGPQSASTAVKERPASSEKVKGG<br>DDHDDTSDSDSDGLTLKELQNRLRRKREQEPTE<br>RPLKGIQSRLRKKRREEGPAETVGSEASDTVEGV<br>LPSKQEPENDQGVVSQAGKDDRESKLEGKAAQD<br>IKDEEPGDLGRPKPECEGYDPNALYCICRQPHNN                                                                                                                                                                                                                                                                                                                          |

SEQ ID Method Predicted Predicted end Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, nucleotide NO: beginning nucleotide location I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, corresponding location T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, to last amino corresponding acid residue of X=Unknown, \*=Stop codon, /=possible nucleotide deletion, to first amino acid residue of peptide \=possible nucleotide insertion peptide sequence sequence RFMICCDRCEEWFHGDCVGISEARGRLLERNGE DYICPNCTILQVQDETHSETADQQEAKWRPGDA **DGTDCTSIGTIEQKSSEDQGIKGRIEKAANPSGKK** KLKIFOPGPGPVPTQLPVLWQVLEIAVSRSISAFT LLHCISCKVIEAPGASKCIGPGCCHVAQPDSVYCS NDCILKHAAATMKFLSSGKEQKPKPKEKMKMK **PEKPSLPKCGAQAGIKISSVHKRPAPEKKETTVK** KAVVVPARSEALGKEAACESSTPSWASDHNYNA VKPEKTAAPSPSLLYKSTKEDRRSEEKAAATAAS KKTAPPGSTVGKQPAPRNLVPKKSSFANVAAAT PAIKKPPSGFKGTIPKRPWLSATPSSGASAARQAG PAPAAATAASKKFPGSAALVGAVRKPVVPSVPM ASPAPGRLGAMSAAPSQPNSQIRQNIRRSLKEIL WK/RFLFFILFRVNDSDDLIMTENEVGKIALHIEK EMFNLFQVTDN/RAYKSKYRSIMFNLKDPKNQG LFHRVLREEISLAKLVRLKPEELVSKELSTWKER PARSVMESRTKLHNESKKTAPRQEAIPDLEDSPP VSDSEEQQESARAVPEKSTAPLLDVFSSMLKDTT SOHRAHLFDLNCKICTGQVPSAEDEPAPKKQKLS ASVKKEDLKSKHDSSAPDPAPDSADEVMPEAVP **EVASEPGLESASHPNVDRTYFPGPPGDGHPEPSPL** EDLSPCPASCGSGVVTTVTVSGRDPRTAPSSSCT AVASAASRPDSTHMVEARQDVPKPVLTSVMVPK SILAKPSSSPDPRYLSVPPSPNISTSESRSPPEGDTT LFLSRLSTIWKGFINMQSVAKFVTKAYPVSGCFD YLSEDLPDTIHIGGRIAPKTVWDYVGKLKSSVSK ELCLIRFHPATEEEEVAYISLYSYFSSRGRFGVVA NNNRHVKDLYLIPLSAQDPVPSKLLPFEGPGKRR LSGWR AQQPVGRRGGASGAGGGRRGTPRPRAGAGPGF 2 3181 3866 OVSSGGCRLSKMRRFLRPGHDPVRERLKRDLFQ FNKTVEHGFPHQPSALGYSPSLRILAIGTRSGAIK LYGAPGVEFMGLHQENNAVTQIHLLPGQCQLVT LLDDNSLHLWSLKVKGGASELQEDESFTLRGPP GAAPSATQITVVLPHSSCELLYLGTESGNVFVVQ LPAFRALEDRTISSDAVLQRLPEEARHRRVFEMV EALOEHPRDPNOILIGYSRGLVVIWDLOGSRVLY HFLSSQQLENIWWQRDGRLLVSCHSDGSYCQWP VSSEAQQPEPLRSLVPYGPFPCKAITRILWLTTRQ G\LPFTIFQGGMPRASYGDRHCISVIHDGQQTAFD FTSRVIGFTVLTEADPAATFDDPYALVVLAEEEL **VVIDLQTAGWPPVQLPYLASLHCSAITCSHHVSN IPLKLWERIIAAGSRQNAHFSTMEWPIDGGTSLTP** APPORDLLLTGHEDGTVRFWDASGVCLRLLYKL STVRVFLTDTDPNENLSAQGEDEWPPLRKVGSF DPYSDDPRLGIQKIFLCKYSGYLAVAGTAGQVLV LELNDEAAEQAVEQVEADLLQDQEGYRWKGHE RLAARSGPVRFEPGFQPFVLVQCQPPAVVTSLAL **HSEWRLVAFGTSHGFGLFDHQQRRQVFVKCTLH** PSDQLALEGPLSRVKSLKKSLRQSFRRMRRSRVS SRKRHPAGPPGEAQEGSAKAERPGLQNMELAPV **ORKIEARSAEDSFTGFVRTLYFADTYLKDSSRHC** PSLWAGTNGGTIYAFSLRVPPAERRMDEPVRAE OAKEIOLMHRAPVVGILVLDGHSVPLPEPLEVAH DLSKSPDMQGSHQLLVVSEEQFKVFTLPKVSAK

Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, SEQ ID Method Predicted Predicted end E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, NO: beginning nucleotide nucleotide location I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, location corresponding corresponding to last amino T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, \*=Stop codon, /=possible nucleotide deletion, acid residue of to first amino acid residue of peptide >=possible nucleotide insertion peptide sequence sequence LKLKLTALEGSRVRRVSVAHFGSRRAEDYGEHH LAVLTNLGDIQVVSLPLLKPQVRYSCIRREDVSGI ASCVFTKYGQGFYLISPSEFERFSLSTKG\LVEPRC LVDSAETKNHRPGNGAGPKKAPSRARNSGTOSD GEEKQPGLVMERALLSDERAATG\VHIEPPWGA **ASAMAEQSEWLSVQAAR** 2 AOOPVGRRGGASGAGGGRRGTPRPRAGAGPGF 3867 3181 QVSSGGCRLSKMRRFLRPGHDPVRERLKRDLFQ FNKTVEHGFPHQPSALGYSPSLRILAIGTRSGAIK LYGAPGVEFMGLHQENNAVTQIHLLPGQCQLVT LLDDNSLHLWSLKVKGGASELQEDESFTLRGPP GAAPSATQITVVLPHSSCELLYLGTESGNVFVVQ LPAFRALEDRTISSDAVLQRLPEEARHRRVFEMV EALQEHPRDPNQILIGYSRGLVVIWDLQGSRVLY **HFLSSQQLENIWWQRDGRLLVSCHSDGSYCQWP** VSSEAQQPEPLRSLVPYGPFPCKAITRILWLTTRO **G\LPFTIFQGGMPRASYGDRHCISVIHDGQQTAFD** FTSRVIGFTVLTEADPAATFDDPYALVVLAEEEL VVIDLQTAGWPPVQLPYLASLHCSAITCSHHVSN **IPLKLWERIIAAGSRQNAHFSTMEWPIDGGTSLTP** APPQRDLLLTGHEDGTVRFWDASGVCLRLLYKL STVRVFLTDTDPNENLSAQGEDEWPPLRKVGSF DPYSDDPRLGIQKIFLCKYSGYLAVAGTAGQVLV LELNDEAAEQAVEQVEADLLQDQEGYRWKGHE RLAARSGPVRFEPGFQPFVLVQCQPPAVVTSLAL HSEWRLVAFGTSHGFGLFDHQQRRQVFVKCTLH **PSDQLALEGPLSRVKSLKKSLRQSFRRMRRSRVS** SRKRHPAGPPGEAQEGSAKAERPGLONMELAPV QRKIEARSAEDSFTGFVRTLYFADTYLKDSSRHC **PSLWAGTNGGTIYAFSLRVPPAERRMDEPVRAE** QAKEIQLMHRAPVVGILVLDGHSVPLPEPLEVAH DLSKSPDMQGSHQLLVVSEEQFKVFTLPKVSAK LKLKLTALEGSRVRRVSVAHFGSRRAEDYGEHH LAVLTNLGDIQVVSLPLLKPQVRYSCIRREDVSGI ASCVFTKYGQGFYLISPSEFERFSLSTKG\LVEPRC LVDSAETKNHRPGNGAGPKKAPSRARNSGTQSD GEEKQPGLVMERALLSDERAATG\VHTEPPWGA ASAMAEQSEWLSVQAAR 2497 3868 Ā **GDSGGPLVCEEPSGRFFLAGIVSWGIGCAEARRP** GVYARVTRLRDWILEATTKASMPLAPTMAPAPA APSTAWPTSPESPVVSTPTKSMQALSTVPLDWVT **VPKLQECGARPAMEKPTRVVGGFGAASGEVPW** QVSLKEGSRHFCGATVVGDRWLLSAAHCFNHT KVEQVRAHLGTASLLGLGGSPVKIGLRRVVLHP LYNPGILDFDLAVLELASPLAFNKYIQPVCLPLAI QKFPVGRKCMISGWGNTQEGNATKPELLOKASV GIIDQKTCSVLYNFSLTDRMICAGFLEGKVDSCQ **VSGIKALYESELADARRVLDETARERARLOIEIG** KLRAELDEVNKSAKKREGELTVAQGRVKDLESL FHRSEVELAAALSDKRGLESDVAELRAQLAKAE DGHAVAKKQLEKETLMRVDLENRCOSLQEELDF RKSVFEEEVRETRRRHERRLVEVDSSRQQEYDFK MAQALEELRSQHDEQVRLYKLELEQTYQAKLDS AKLSSDQNDKAASAAREELKEARMRLESLSYQL SGLQKQASAAEDRIRELEEAMAGERDKFRKMLD

|               |        |                                                                      |                                                                                              | A Alaria Caratia Danasia And                                                                                                                                                                                                                                                                                                                 |
|---------------|--------|----------------------------------------------------------------------|----------------------------------------------------------------------------------------------|----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| SEQ ID<br>NO: | Method | Predicted beginning nucleotide location corresponding to first amino | Predicted end<br>nucleotide<br>location<br>corresponding<br>to last amino<br>acid residue of | Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, |
|               |        | acid residue of peptide sequence                                     | peptide<br>sequence                                                                          | \=possible nucleotide insertion                                                                                                                                                                                                                                                                                                              |
|               |        |                                                                      |                                                                                              | AKEQEMTEMRDVMQQQLAEYQELLDVKLALD<br>MEINAYRKLLEGEEERLKLSPSPSSRVTVSRATSS                                                                                                                                                                                                                                                                       |
|               |        |                                                                      |                                                                                              | SSGSLSATGRLGRSKRKR\WRWRSPW\QRPKRPG<br>HGHGWQRWLPPGPAGLGLGQR\HIEEIDLEGKFV                                                                                                                                                                                                                                                                     |
|               |        |                                                                      |                                                                                              | QLKNNSDKDQSLGNWRIKRQVLEGEEIAYKFTP<br>KYILRAGQMVTVWAAGAGVAHSPPSTLVWKGQ                                                                                                                                                                                                                                                                        |
|               | }      |                                                                      |                                                                                              | SSWGTGESFRTVLVNADGEEVAMRTVKKSSVM<br>RENENGEEEEEEAEFGEEDLFHQQGDPRTTSRGC                                                                                                                                                                                                                                                                       |
|               |        |                                                                      |                                                                                              | YVM                                                                                                                                                                                                                                                                                                                                          |
| 3869          | A      | 1                                                                    | 1942                                                                                         | RYRAGIPGDGRKDYIRLTRPGLTLPGRAMFARGS<br>RRRRSGRAPPEAEDPDRGQPCNSCREQCPGFLLH                                                                                                                                                                                                                                                                     |
| !             |        |                                                                      |                                                                                              | GWRKICQHCKCPREEHAVHAVPVDLERIMCRLIS                                                                                                                                                                                                                                                                                                           |
|               |        |                                                                      |                                                                                              | DFQRHSISDDDSGCASEEYAWVPPGLKPEQVYQ<br>FFSCLPEDKVPYVNSPGEKYRIKQLLHQLPPHDS                                                                                                                                                                                                                                                                      |
|               | }      | 1                                                                    | }                                                                                            | EAQYCTAL\EE\EEKKELRAFSQQRKRENLG/RLG                                                                                                                                                                                                                                                                                                          |
|               |        |                                                                      |                                                                                              | IVRIFPVTIT\GAI\CEECGKQIGGGDIAVF\ASRASL                                                                                                                                                                                                                                                                                                       |
|               |        |                                                                      |                                                                                              | GLLLGQPSCF\VCTTCQELLVDLIYFYHVGKVYC<br>GRHHAECLRPRCQACDEIIFSPECTEAEGRHWHM                                                                                                                                                                                                                                                                     |
|               |        |                                                                      |                                                                                              | DHFCCFECEASLGGQRYVMRQSRPHCCACYEAR                                                                                                                                                                                                                                                                                                            |
|               | 1      |                                                                      |                                                                                              | HAEYCDGCGEHIGLDQGQMAYEGQHWHASDRC                                                                                                                                                                                                                                                                                                             |
| •             |        |                                                                      |                                                                                              | FCCSRCGRALLGRPFLPRRGLIFCSRACSLGSEPT<br>APGPSRRSWSAGPVTAPLAASTASFSAVKGASET                                                                                                                                                                                                                                                                    |
|               |        | 1                                                                    |                                                                                              | TTKGTSTELAPATGPEEPSRFLRGAPHRHSMPEL                                                                                                                                                                                                                                                                                                           |
|               |        |                                                                      |                                                                                              | GLRSVPEPPPESPGQPNLRPDDSAFGRQSTPRVSF                                                                                                                                                                                                                                                                                                          |
|               |        |                                                                      |                                                                                              | RDPLVSEGGPRRTLSAPPAQRRRPRSPPPRAPSRR<br>RHHHHNHHHHHNRHPSRRRHYQCDAGSGSDSE                                                                                                                                                                                                                                                                      |
|               |        |                                                                      | *                                                                                            | SCSSSPSSSSSESSEDDGFFLGERIPLPPHLCRPMP                                                                                                                                                                                                                                                                                                         |
|               |        |                                                                      |                                                                                              | AQDTAMETFNSPSLSLPRDSRAGMPRQARDKNC<br>IVA                                                                                                                                                                                                                                                                                                     |
| 3870          | A      | 2                                                                    | 3485                                                                                         | FVWRVFYVHASCMPPRARSWEGAHAPVGMHV<br>AEAHACSSQQQQMPPAQFWMLEWLLHLCAFLS                                                                                                                                                                                                                                                                          |
|               |        | 1                                                                    |                                                                                              | TPSFPHWCCCSNPHGSIADKPEEIVPASKPSRAAE                                                                                                                                                                                                                                                                                                          |
|               | İ      |                                                                      |                                                                                              | NMAVEPRVATIKQRPSSRCFPAGSDMNSVYERQ                                                                                                                                                                                                                                                                                                            |
|               |        |                                                                      |                                                                                              | GIAVMTPTVPGSPKAPFLGIPRGTMRRQKSIDSRI<br>FLSGITEEERQFLAPPMLKFTRSLSMPDTSEDIPPP                                                                                                                                                                                                                                                                  |
|               |        |                                                                      |                                                                                              | POSVPPSPPPPSPTTYNCPKSPTPRVYGTIKPAFNQ                                                                                                                                                                                                                                                                                                         |
|               | 1      |                                                                      | 1                                                                                            | NSAAKVSPATRSDTVATMMREKGMYFRRELDR                                                                                                                                                                                                                                                                                                             |
|               |        |                                                                      | ļ                                                                                            | YSLDSEDLYSRNAGPQANFRNKRGQMPENPYSE<br>VGKIASKAVYVPAKPARRKGMLVKQSNVEDSPE                                                                                                                                                                                                                                                                       |
|               |        |                                                                      |                                                                                              | KTCSIPIPTIIVKEPSTSSSGKSSQGSSMEIDPQAPE                                                                                                                                                                                                                                                                                                        |
|               |        | 1                                                                    |                                                                                              | PPSQLRPDESLTVSSPFAAAIAGAVRDREKRLEA                                                                                                                                                                                                                                                                                                           |
|               | 1.     | }                                                                    | 1                                                                                            | RRNSPAFLSADLGDEHVGLGPPAPRTRPSMFPEE<br>GDFADEDSAEQLSSPMPSATPREPENHFVGGAEA                                                                                                                                                                                                                                                                     |
|               |        |                                                                      |                                                                                              | SAPGEAGRPLNSTSKAQGPESSPAVPSASSGTAG                                                                                                                                                                                                                                                                                                           |
|               |        |                                                                      | j                                                                                            | PGNYVHPLTGRLLDPSSPLALALSARDRAMKES                                                                                                                                                                                                                                                                                                            |
|               |        |                                                                      |                                                                                              | QQGPKGEAPKADLNKPLYIDTKMRPSLDAGFPT<br>VTRQNTRGPLRRQETENKYETDLGRDRKGDDK                                                                                                                                                                                                                                                                        |
|               | 1      |                                                                      |                                                                                              | KNMLIDIMDTSQQKSAGLLMVHTVDATKLDNA                                                                                                                                                                                                                                                                                                             |
|               |        |                                                                      |                                                                                              | LQEEDEKAEVEMKPDSSPSEVPEGVSETEGALQI                                                                                                                                                                                                                                                                                                           |
|               | 1      |                                                                      |                                                                                              | SAAPEPTTVPGRTIVAVGSMEEAVILPFRIPPPPLA<br>SVDLDEDFIFTEPLPPPLEFANSFDIPDDRAASVPA                                                                                                                                                                                                                                                                 |
|               |        |                                                                      |                                                                                              | LSDLVKQKKSDTPQSPSLNSSQPTNSADSKKPAS                                                                                                                                                                                                                                                                                                           |
|               |        |                                                                      |                                                                                              | LSNCLPASFLPPPESFDAVADSGIEEVDSRSSSDH                                                                                                                                                                                                                                                                                                          |
|               |        |                                                                      | ]                                                                                            | HLETTSTISTVSSISTLSSEGGENVDTCTVYADGQ<br>AFMVDKPPVPPKPKMKPIIHKSNALYQDALVEE                                                                                                                                                                                                                                                                     |
|               |        | 1                                                                    |                                                                                              |                                                                                                                                                                                                                                                                                                                                              |

| SEQ ID<br>NO: | Method | Predicted beginning nucleotide location corresponding to first amino acid residue of peptide sequence | Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence | Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion                          |
|---------------|--------|-------------------------------------------------------------------------------------------------------|------------------------------------------------------------------------------------------------|-------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
|               |        |                                                                                                       |                                                                                                | DVDSFVIPPPAPPPPPGSAQPGMAKVLQPRTSKL WGDVTEIKSPILSGPKANVISELNSILQQMNREKL AKPGEGLDSPMGAKSASLAPRSPEIMSTISGTRST TVTFTVRPGTSQPITLQSRPPDYESRTSGTRRAPS PVVSPTEMNKETLPAPLSAATASPSPALSDVFSLP SQPPSGDLFGLNPAGRSRSPSPSILQQPISNKPFTT KPVHLWTKPDVADWLESLNLGEHKEAFMDNEI DGSHLPNLQKEDLIDLGVTRVGHRMNIERALKQ LLDR                                                                                                       |
| 3871          | A      | 35                                                                                                    |                                                                                                | VESRSAWHEGEDQIDRLDFIRNQMNLLTLDVKK KIKEVTEEVANKVSCAMTDEICRLSVLVDEFCSE FHPNPDVLKIYKSELNKHIEDGMGRNLADRCTD EVNALVLQTQQEIIENLKPLLPAGIQDKLHTLIPC KKFDLSYNLNYHKLCSDFQEDIVFRFSLGWSSLV HRFLGPRNAQRVLLGLSEPIFQLPRSLASTPTAPT TPATPDNASQEELMITLVTGLASVTSRTSMGIIIV GGVIWKTIGWKLLSVSLTMYGALYLYERLSWTT HAKERAFKQQFVNYATEKLRMIVSSTSANCSHQ VKQQIATTFARLCQQVDITQKQLEEIARLPKEID QLEKIQNNSKLLRNKAVQLENELENFTKQFLPSS NEES  |
| 3872          | A      | 35                                                                                                    |                                                                                                | VESRSAWHEGEDQIDRLDFIRNQMNLLTLDVKK KIKEVTEEVANKVSCAMTDEICRLSVLVDEFCSE FHPNPDVLKIYKSELNKHIEDGMGRNLADRCTD EVNALVLQTQQEIIENLKPLLPAGIQDKLHTLIPC KKFDLSYNLNYHKLCSDFQEDIVFRFSLGWSSLV HRFLGPRNAQRVLLGLSEPIFQLPRSLASTPTAPT TPATPDNASQEELMITLVTGLASVTSRTSMGIIIV GGVIWKTIGWKLLSVSLTMYGALYLYERLSWTT HAKERAFKQQFVNYATEKLRMIVSSTSANCSHQ VKQQIATTFARLCQQVDITQKQLEEEIARLPKEID QLEKIQNNSKLLRNKAVQLENELENFTKQFLPSS NEES |
| 3873          | Α      | 2944                                                                                                  | 2089                                                                                           | PVCTALTPGRMTDDKDVLRDVWFGRIPTCFTLY QDEITEREAEPYYLLLPRVSYLTLVTDKVKKHFQ KVMRQEDISEIWFEYEGTPLKWHYPIGLLFDLLA SSSALPWNITVHFKSFPEKDLLHCPSKDAIEAHF MSCMKEADALKHKSQVINEMQKKDHKQLWMG LQNDRFDQFWAINRKLMEYPAEENGFRYIPFRIY QTTTERPFIQKLFRPVAADGQLHTLGDLLKEVCP SAIDPEDGEKKNQVMIHGIEPMLETPLQWLSEHL SYPDNFLHISIIPQPTD                                                                                                 |
| 3874          | A      | 776                                                                                                   | 366                                                                                            | QARGAPSSPMCPLPLAAAAVAAPRAPLRLLNRG<br>LAAAMSTAQSLKSVDYEVFGRVQGVCFRMYTE<br>DEARKIGVVGWVKNTSKGTVTGQVQGPEDKVN<br>SMKSWLSKVGSPSSRIDRTNFSNEKTISKLEYSNF<br>SIRY                                                                                                                                                                                                                                              |
| 3875          | A      | 1081                                                                                                  | 182                                                                                            | SLSSCQTDPRPMSAPLDAALHALQEEQARLKMR LWDLQQLRKELGDSPKDKVPFSVPKIPLVFRGHT QQDPEVPKSLVSNLRIHCPLLAGSALITFDDPKVA EQVLQQKEHTINMEECRLRVQVQPLELPMVTTIQ VMVSSQLSGRRVLVTGFPASLRLSEEELLDKLEIF FGKTRNGGGDVDVRELLPGSVMLGFARDGVAQ RLCQIGQFTVPLGGQQVPLRVSPYVNGEIQKAEI RSQPVPRSVLVLNIPDILDGPELHDVLEIHFQKPT                                                                                                               |

| SEQ ID<br>NO: | Method  | Predicted beginning nucleotide location corresponding to first amino acid residue of peptide sequence | Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence | Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion                                                                              |
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|               |         |                                                                                                       |                                                                                                | RGGGEVEALTVVPQGQQGLAVFTSESG                                                                                                                                                                                                                                                                                                                                                                                                                               |
| 3876          | A       | 26                                                                                                    | 431                                                                                            | RMMKCPQALLAIFWLLLSWVSSEDKVVQSPLSL<br>VVHEGDTVTLNCSYEVTNFRSLLWYKQEKKAPT<br>FLFMLTSSGIEKKSGRLSSILDKKELSSILNITATQ<br>TGDSAIYLCAVEAQCSLVTCSLYSNSTAEALQL                                                                                                                                                                                                                                                                                                       |
| 3877          | A       | 3                                                                                                     |                                                                                                | KAFRLLAERGAAAAMLWSGCRRFGARLGCLPG GLRVLVQTGHRSLTSCIDPSMGLNEEQKEFQKV AFDFAAREMAPNMAEWDQKELFPVDVMRKAA QLGFGGVYIQTDVGGSGLSRLDTSVIFEALATGC TSTTAYISIHNMCAWMIDSFGNEEQRHKFCPPLC TMEKFASYCLTEPGSGSDAASLLTSAKKQGDHYI LNGSKAFISGAGESDIYVVMCRTGGPGPKGISCIV VEKGTPGLSFGKKEKKVGWNSQPTRAVIFEDCA VPVANRIGSEGQGFLIAVRGLNGGRINIASCSLGA AHASVILTRDHLNVRKQFGEPLASNQYLQFTLA DMATRLVAARLMVRNAAVALQEERKDAVALCS MAKLFATDECFAICNQALQMHGGYGYLKDYAV QQYVRDSRVHQILEGSNEVMRILISRSLLQE |
| 3878          | <b></b> | 10                                                                                                    | 1014                                                                                           | LPGSTISSSGCQAPGRADSSGGARNSRRGDSRPG                                                                                                                                                                                                                                                                                                                                                                                                                        |
| 3078          | A       |                                                                                                       | 1014                                                                                           | SCNRQAVAPPCPSPGPQSRHWIHRGTAPQAGETR TLGRGSSAPNACSASVTPCCPSSPPS*SCL*PTRRS PQNSSSTEVYRGFWQHGLPST**PFSS*QWPGQH TQGCSKLLGKQTTHLPCSTWPA**PSPSCLTRFR* W*PSLMCLWASSCSVCV*SPSGSCRH*LWGTHST SRTC*ARRSSALPTGLCTDDTSWASSSKARPCAL QRPSSLSSLSPCLTC*W*LSSSSPMSARSPAGAET GSWATGSPRLTQWKSSRLTSTSHSARSAWKPSA TESTPSWPRFSSWTSGEDPASPAPAI                                                                                                                                     |
| 3879          | A       | 200                                                                                                   | 699                                                                                            | LLLTGYIQTLQNQQLSGNQQEMQAVDNLTSAPG NTSLCTRDYKITQVLFPLLYTVLFFVGLITNGLA MRIFFQIRSKSNFIIFLKNTVISDLLMILTFPFKILS DAKLGTGPLRTFVCQVTSVIFYFTMYISISFLGLIT IDRYQKTTRPFKTSNPKNLLGAKILK                                                                                                                                                                                                                                                                                |
| 3880          | A       | 26                                                                                                    | 169                                                                                            | QPETDTMVHLTPEEKSAVTALWGKVNVDEDAG<br>DDLCQILVDRPRLRI                                                                                                                                                                                                                                                                                                                                                                                                       |
| 3881          | A       | 37                                                                                                    | 1100                                                                                           | TPLFDFWPGFVLSWLQPLSASLRARRAASGPPAC RIMPTTVDDVLEHGGEFHFFQKQMFFLLALLSAT FAPIYVGIVFLGFTPDHRCRSPGVAELSLRCGWSP AEELNYTVPGPGPAGEASPRQCRRYEVDWNQST FDCVDPLASLDTNRSRLPLGPCRDGWVYETPGSS IVTEFNLVCANSWMLDLFQSSVNVGFFIGSMSIG YIADRFGRKLCLLTTVLINAAAGVLMAISPTYTW MLIFRLIQGLVSKAGWLIGYILITEFVGRRYRRTV GIFYQVAYTVGLLVLAGVAYALPHWRWLQFTV ALPNFFFLLYYWCIPESPRWLISQNKNAEAMRIIK HIAKKNGKSLPASL                                                                              |
| 3882          | A .     | 573                                                                                                   |                                                                                                | KSKCRFPEGLSEGFGPMRKEALSSGSVQEAEAM LDEPQEQAEGSLTVYVISEHSSLLPQDMMSYIGP KRTAVVRGIMHREAFNIIGRRIVQVAQAMSLTED VLAAALADHLPEDKWSAEKRRPLKSSLGYEITFS LLNPDPKSHDVYWDIEGAVRRYVQPFLNALGAA GNFSVDSQILYYAMLGVNPRFDSASSSYYLDMH SLPHVINPVESRLGSSAASLYPVLNFLLYVPELAH SPLYIQDKDGAPVATNAFHSPRWGGIMVYNVDS KTYNASVLPVRVEVDMVRVMEVFLAQLRLLFGI                                                                                                                                    |

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|               |        |                                                                                                       |                                                                                                | AQPQLPPKCLLSGPTSEGLMTWELDRLLWARSV<br>ENLATATTTLTSLA                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          |
| 3883          | A      | 2369                                                                                                  | 844                                                                                            | RIHREEDFQFILKGIARLLSNPLLQTYLPNSTKKIQ FHQELLVLFWKLCDFNKVGQPRGALQGDGEQLP Q*PGGRDSVRLRGVGQSCPSLELSPLGPSPHP*KF LFFVLKSSDVLDILVPILFFLNDARADQSRVGLM HIGVFILLLLSGECNFGVRLNKPYSIRVPMDIPVF TGTHADLLIV\VFHKIITSGHQRLQPLFDCLLTIVV NVSPYLKSLSMVTANKLLHLLEAFSTTWFLFSAA QNHHLVFFLLEVFNNIIQYQFDGNSNLVYAIIRKR SIFHQLANLPTDPPTIHKALQRRRRTPEPLSRTGS QGGAPPWRAPAPLPLQSQAPSRPVWWLLQALTS *PRSPRCQRMAPCGPWNLSPSRAWRMAARLRGS PARHGGSSGDRP/HSSASGQWSPTPEWVLSWKS KLPLQTIMRLLQVLVPQVEKICIDKGLTDESEILR FLQHGTLVGLLPVPHPILIRKYQANSGTAMWFRT YMWGVIYLRNVDPPVWYDTDVKLFEIQRV |
| 3884          | A      | 1                                                                                                     | 804                                                                                            | NGPRAPFSQEGQSTGPPPLIPRLGQHGAQGRIPPL NPGQGPGPNKDDSRGPPNHHMGPMSERRHEQSG GPEHGPERGPLRGGQDCRGPPDRRGPHPDFPDDF SRPDDFHPDKRFGHRLREFEGRGGPLPQEEKWR RGGPGPPFPPDHREFSEGDGRGAARGPPGAWEG RRPGG*TFPPGSRGPTFS/SGAEEESFRRGAPPRHE GRAPPRGRDGFPGPEDFGPEENFDASEEAARGRD LRGRGRGTPRGERVTKDTWSGRIGCRIHWL                                                                                                                                                                                                                                                          |
| 3885          | A      | 3                                                                                                     | 996                                                                                            | GRRRAGPAHSARMYNMMETELKPPGPQQTSGG GGGNSTAAAAGGNQKNSPDRVKRPMNAFMVW SRGQRRKMAQENPKMHNSEISKRLGAEWKLLSE TEKRPFIDEAKRLRALHMKEHPDYKYRPRRKTK TLMKKDKYTLPGGLLAPGGNSMASGVGVGAGL GAGVNQRMDSYAHMNGWSNGSYSMMQDQLG YPQHPGLNAHGAAQMQPMHRYDVSALQYNSM TSSQTYMNG/SRPTYSMSYSQQGTPGMAPGS\MG SVVKSEASSSPPVVTSSSHSRAPCQAGDLRDMIS MYLPGAEVPEPAAPSRLHMSQHYQSGPVPGTAI NGTLPLSHM                                                                                                                                                                                       |
| 3886          | A      | 773                                                                                                   | 317                                                                                            | QCTQKAAEGYTQFYYVDVLDGKLACVNKCTKG<br>TKSQMNCNLGTCQLQRSGPRCLCPNTNTHWYW<br>GETCEFNIAKSLVYGIVGAVMAVLLLALIILIILFS<br>LSQ\RKRHRPESEGEADFGLENATNNFG\PTLETV<br>DSGTELHIQ\RPEMVASTV                                                                                                                                                                                                                                                                                                                                                                   |
| 3887          | A      | 3                                                                                                     | 466                                                                                            | VDFRVKTLLVDNKCFVLQLWDTAGQERYHSMT<br>RQLLRKADGVVLMYDITSQESFAHVRYWLDCL<br>QDAGSDGVVILLLGNKMDCEEERQVSVEAGQQL<br>AQELGVYFGECSAALGHNILEPVVNLARSLRMQ<br>EEGLKDSLVKVAPKRPPKRFGCCS                                                                                                                                                                                                                                                                                                                                                                   |
| 3888          | A      | 3412                                                                                                  | 3144                                                                                           | QNIDITNFSSSWNDGLAFCALLHTYLPAHIPYQEL<br>NSQDKRRNFMLAFQAAESVGIKSTLDINEMVRT<br>ERPDWQNVMLYVTAIYKYFET                                                                                                                                                                                                                                                                                                                                                                                                                                            |
| 3889          | A      | 1                                                                                                     | 1160                                                                                           | LVVTAITAILAFPNEYTRMSTSELISELFNDCGLL DSSKLCDYENRFNTSKGGELPDRPAGVGVYSAM WQLALTLILKIVITIFTFGMKIPSGLFIPSMAVGAI AGRLLGVGMEQLAYYHQEWTVFNSWCSQGAD CITPGLYAMVGAAACLGGVTRMTVSLVVIMFEL TGGLEYIVPLMAAAMTSKWVADALGREGIYDA                                                                                                                                                                                                                                                                                                                                |

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| ·             |        |                                                                                                       |                                                                                                | HIRLNGYPFLEAKEEFAHKTLAMDVMKPRRNDP<br>LLTVLTQDSMTVEDVETIISETTYSGFPVVVSRES<br>QRLVGFVLRRDLIISIENARKKQDGVVSTSIIYFTE<br>HSPPLPPYTPPTLKLRNILDLSPFTVTDLTPMEIVV<br>DIFRKLGLRQCLVTHNGRLLGIITKKDVLKHIAQ<br>MANQDPDSILFN                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 |
| 3890          | A      | 1                                                                                                     | 387                                                                                            | SWCWTGIFVLGTTNLRLEGSWYRSLWGPGFNTT<br>TATLGFGAPQAPVGDVALNQPDMCVYRRGRKK<br>RVPYTKLQLKELENEYAINKFINKDKRRRISAAT<br>NLSERQVTIWFQNRRVKDKKIVSKLKDTVS                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  |
| 3891          | A      | 2                                                                                                     | 2914                                                                                           | RGGGGDHKMADLSLLQEDLQEDADGFGVDDYS SESDVIIIPSALDLAST/QDEMVERPLGRL\DK\YA ASENHI*PDKMVAPEFASIPLRE\VCDDERDCIAV LGKN*PDWADDSEPT\VRAAELEQVPHIALFLFK KTRLSITICFFSKFLLPYCGLDTLADQN\NQVRKT SQAALL\ALLEQELIERFDVETKVCPVLIELTAPDS NDDVKTEAVAIMCKMAP\MVGKDITERLILPRFC EMCCDCRMFH\VRK\VCAANFGDICSVVGQQAT EEMLLPRFFQLCSDNVWGVRKACAECFMAVSC ATCQEIRRTKLSALFINLISDPSRWVRQAAFQSLG PFISTFANPSSSGQYFKEESKSSEEMSVENNKRTR DQEAPEDVQVRPEDTPSDLSVSNSSVILENTMED HAAEASGKPLGEISVPLDSSLLCTLSSESHQEAAS NENDKKPGNYKSMLRPEVGTTSQDSALLDQELY NSFHFWRTPLPEIDLDIELEQNSGGKPSPEGPEEE SEGPVPSSPNITMATRKELEEMIENLEPHIDDPDV KAQVEVLSAALRASSLDAHEETISIEKRSDLQDE LDINELPNCKINQEDSVPLISDAVENMDSTLHYIH NDSDLSNNSSFSPDEERRTKVQDVVPQALLDQY LSMTDPSRAQTVDTEIAKHCAYSLPGVALTLGR QNWHCLRETYETLASDMQWKVRRTLAFSIHELA VILGD\QLTAADLVPIFNGFLK*PSMKSRIGVLKH LHDFLKLLHIDKRREYLYQLQEFLVTDNSRNWR FRAELAEQLILLLELYSPRDVYDYLRPIALNLCAD KVSSVRWISYKLVSEMVKKLHAATPPTFGVDLIN ELVENFGRCPKWSGRQAFVFVCQTVIEDDCLPM DQFAVHLMPHLLTLANDRVPNVRVLLAKTLRQT LLEKDYFLASASCHQEAVEQTIMALQMDRDSDV KYFASIHPASTKISEDAMSTASSTY |
| 3892          | A      | 158                                                                                                   | 2191                                                                                           | VPLPAPSGLSGGGSRGAGCKKAPPGRAPAPGLAP LRPSEPTMAVPPGHGPFSGFPGPQEHTQVLPDVR LLPRRLPLAFRDATSAPLRKLSVDLIKTYKHINEV YYAKKKRRAQQAPPQDSSNKKEKKVLNHGYDD DNHDYIVRSGERWLERYEIDSLIGKGSFGQVVKA YDHQTQELVAIKIIKNKKAFLNQAQIELRLLELM NQHDTEMKYYIVHLKRHFMFRN\HLCLVFELLS YNLYDLLRNTHFRGVSLNLTRKLAQQLCTALLF LATPELSIIHCDLKPENILLCNPKRSAIKIVDFGSS CQLGQRIYQYIQSRFYRSPEVLLGTPYDLAIDMW SLGCILVEMHTGEPLFSGSNEVCPQEGVDQMNRI VEVLGIPPAAMLDQAPKARKYFERLPGGGWTLR RTKELRKDYQGPGTRRLQEVLGVQTGGPGGRRA GEPGHSPAD\Y\LRFQDLVLRMLEYEPAARISPLG ALQHGFFRRTADEATNTGPAGSSASTSPAPLDTC PSSSTASSISSSGGSSGSSGSSDNRTYRYSNRYCGGP                                                                                                                                                                                                                                                                                                                                                                                                                                                               |

| Method   Predicted   beginning   nucleated   beginning   nucleated   beginning   nuclear   beginning   to first amino   sold are alton   beginning   to first amino   sold are alton   beginning   to first amino   sold are alton   beginning   beg   |               |        |                                                                          |                                                                          |                                                                                                                                                                                                                                                        |
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| RMTGGRPPLPPPDDATLGPHLGLRGVPQSTAAS   S   S                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      |               |        |                                                                          |                                                                          | QAPASASSLPGTGAQLPPQPRYLGRPPSPTSPPPP                                                                                                                                                                                                                    |
| 3893   A   68   258                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            |               | !      |                                                                          |                                                                          | RMTGGRPPLPPPDDPATLGPHLGLRGVPQSTAAS                                                                                                                                                                                                                     |
| 3894 A                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         | 3893          | A      | 68                                                                       | 258                                                                      |                                                                                                                                                                                                                                                        |
| AASGSHPEVGSVLQRSSQPHWPNPWGAGHLPP PAGPFPYNPPAGPGAAAGL*SPPRSSPTPCSVOP QSCRANASAPPAGPCLAGAPPAASLPPGFOSVS AAPAPGGPAPABPLGVPPVPAWILIPDSPPLPGT HSGPPPAAVSLPPAAAACPVVVPPPLPHPDLES PSAAAPNPGCAGGIRHFPPGSPEASSPLRPAAAPA LLPLPRPS*PVPWKPLHSPVAVAGGSFVAGGSV LPAPDLDQPRPSGPPAASPTTGPGVAVAGGSFVAGGSV LPAPDLDQPRPSGPPAASPTTGPGVAGGSFVAGGSV LPAPDLDQPRPSGPPAASPTTGPGVAGGSFVAGGSV LPAPDLDQPRPSGPPAASPTTGPGVAGGSFVAGGSV LPAPDLDQPRPSGPPAASPTTGPGVAGGSFVAGGSV LPAPDLDQPRPSGPPAASPTTGPGVAGGSFVAGGSV LPAPDLDQPRPSGPPAASPTTGPGVAGGSFVAGGSV LPAPDLDQPRPSGPPAASPTTGPGVAGAGATRW PLSCSIDFQRLLAHEETQKRRAKESGMAFTQLI FRDVALEFSQDEWKCLINSTQRTLYRDVMLENYR NLVSLDLSRNCVIKELAPQGENP/ARSIPHSDIGT T*KT*H*RVLLQGNQEKNTRL*LSVER**KKLQQ SDYGPKRRSYL*GERTK*KRYRQVYTSAVLSTSAVLSF PQIISSTIKTHVSNK YGTDFICSSLLTQEQKSCIRE KPYXYECDKALNHGSHMTVRQVSHSSOKSIRE KPYXYECDKALNHGSHMTVRQVSHSGSKOYKC DLCGKVFSQKSNLARHWRVHTGEKPYKCNECO RSFSRNSCLALHRV*HTGEKPYKCNECO RSFSRNSCLALHRV*HTGEKPYKCNECO RSFSRNSCLALHRV*HTGEKPYKCNECO RSFSRNSCLALHRV*HTGEKPYKCNECO RSFSRNSCLALHRV*HTGEKPYKCNECO RSFSRNSCLALHRV*HTGEKPYKCNECO RSFSRNSCLALHRV*HTGEKPYKCNECO RSFSRNSCLALHRV*HTGEKPYKCNECO RSFSRNSCLALHRV*HTGEKPYKCNECO RSFSRNSCLALHRV*HTGEKPYKCNECO RSFSRNSCLALHRV*HTGEKPYKCNECO RSFSRNSCLALHRV*HTGEKPYKCNECO RSFSRNSCLALHRV*HTGEKPYKCNECO RSFSRNSCLALHRV*HTGEKPYKCNECO RSFSRNSCLALHRV*HTGEKPYKCNECO RSFSRNSCLALHRV*HTGEKPYKCNECO RSFSRNSCLALHRV*HTGEKPYKCNECO RSFSRNSCLALHRV*HTGEKPYKCNECO RSFSRNSCLALHRV*HTGEKPYKCNECO RSFSRNSCLALHRV*HTGEKPYKCNECO RSFSRNSCLALHRV*HTGEKPYKCNECO RSFSRNSCLALHRV*HTGEKPYKCNECO RSFSRNSCLALHRV*HTGEKPYKCNECO RSFSRNSCLALHRV*HTGEKPYKCNECO RSFSRNSCLALHRV*HTGEKPYKCNECO RSFSRNSCLALHRV*HTGEKPYKCNECO RSFSRNSCLALHRV*HTGEKPYKCNECO RSFSRNSCLALHRV*HTGEKPYKCNECO RSFSRNSCLALHRV*HTGEKPYKCNECO RSFSRNSCLALHRV*HTGEKPYKCNECO RSFSRNSCLALHRV*HTGEKPYKCNECO RSFSRNSCLALHRV*HTGEKPYKCNECO RSFSRNSCLALHRV*HTGEKPYKCNECO RSFSRNSCLALHRV*HTGEKPYKCNECO RSFSRNSCLALHRV*HTGEKPYKCNECO RSFSRNSCLALHRV*HTGEKPYKCNECO RSFSRNSCLALHRV*HTGEKPYKCNECO RSFSRNSCLALHRV*HTGEKPYKCNECO RSFSRNSCLALHRV*HTGEKPYKCNECO RSFSRNSCLALHRV*HTGEKPYKCNECO RSFSRNSCLALHRV*HTGEKPYKCNECO RSFSR | 3894          | Α      | 1120                                                                     | 136                                                                      |                                                                                                                                                                                                                                                        |
| LPAPDLDQPPPSGPPAASPTPGPGVAQPPPGSAVL                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            |               |        | 1120                                                                     |                                                                          | AASGSHPEVGSVLQRSSQPHWPNPWPGAGHLPP<br>PAGPFPYNPPAGPGAAAGLA*SPPRSSPTPCSVGP<br>QSCPANASAPPAQPCLAGAPPAASLPPPGPGSVS<br>AAPAPGGPAPAEPPLGVPPVPAWLLPDSPPLPGT<br>HSGPPPAAVSLPPAAAACPVVVPPPLPHHPPDLES<br>PSAAAPNPGCAGGIRHFPPGSPEASSPLRPAAAPA                     |
| SLCSSQACVLPVPGPSLLLPQGLHVGCASAGTRW PLSCSIDFQRLLAHEETQKRRAKESGMAFTQLI FRDVALEFSQDEWKCLNSTQRTLYRDVMLENYR NLVSLDLSRNCVIKELAPQQEGNP/ARSIPHSDIGT T*KT*H*RV1LQGNQEKNTRL*LSVER**KKLQQ SDYGPKRKSYL*ERPTR*KRYRKQVY*TSA\*LSF LPHPHELQQFQAEGKIYECNHVEKSVNHGSSVSP PQIISSTIKTHVSNKYGTDFICSSLLTQEQKSCIRE KPYRYIECDKALNHGSHMTVRQVSHSGEKGYKC DLCGKVFSQKSNLARHWRVHTGEKPYKCNECD RSFSRNSCLALHRRVHTGEKPYKCYECDKVFSR NSCLALHQKTHIGEKPYTCKECGQAFSVRSTLTN HQVIHSDK  3896 A 202 498 MVQSCSAYGCKNRYDKDKPVSFHKFPLTRPSLC KEWEAAVRRKNFKPTKYSSICSEHFTPDCFKREC NNKLLKENAVPTITLCTBPHDKKEDLLEPQEQ 3897 A 2 382 SHGLSRAPHLSAAPAPALASRPCFSSAPCSQGGG GGGPATMIHFILLFSRQGKLRLQKWYITLPDKER KKITREIVQIILSRGHRTSSFVDWKELKLVYKRYA SLYFCCAIENQDNELLTLENVHR 3898 A 718 305 SEQEPLLGDTPGSREWDILETBEHYKSRWRSIRIL YLTMFLSSVGFSVVMMSIWPYLQKIDPTADTSFL GWVIASYSLGQMVASPIFGLWSNYRPRKEPLIVSI LISVAANCLYAYLHIPASHNKYYMLVARGLLGIG GAMAERKGTAKVDFLKKIEKEIQQKWDTERVFE VNASNLEKQTSKGKYFVTFPYPYMNGRLHLGHT FSLSKCEFAVGYQRLKGKCCLPFFGLHCTGMPIK ACADKLKREIELY/GCPPDFPDEEEEEEETSVKTE DIIIKDKAKGKKSKAA/AKAGSSKYQWGIMKSLG LSDEEIVKFSEAEHWLDYFNALAIQDLKRMG 3900 A 360 1 VPATSSNVSPSSSESSEPDLSSRSSSSDAPSSSPSVP SPCSLSLSSPESPLLPTLLSSKSPAGSAGPTCGCPS GPGLRATA/PSRLSSSIAAH/SSSAPETSRPAAARE RSPPLHDRESHE 3901 A 193 345 GEWAVPFAPGGQGVSIPHGPEPGQGSGVHIAPRQ                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        |               |        | ·                                                                        |                                                                          | LPAPDLDQPRPSGPPAASPTPGPGVAQPPPGSAVL                                                                                                                                                                                                                    |
| SLCSSQACVLPVPGPSLLLPQGLHVGCASAGTRW PLSCSIDFQRLLAHEETQKRRAKESGMAFTQLT FRDVALEFSQDEWKCLNSTQRTLYRDVMLENYR NLVSLDLSRNCVIKELAPQQEGNP/ARSIPHSDIGT T*KT*H*RVLLQGNQEKNTRL*LSVER**KKLQQ SDYGPKKSYL*ERPTR*KRYRKQVY*TSA\*LSF LPHPHELQQFQAEGKLYECNHVEKSVNHGSSVSP PQIISSTIKTHVSNKYGTDFICSSLLTQEQKSCIRE KPYRYIECDKALNHGSHMTVRQVSHIGEKGYKC DLCGKVFSQKSSNLARHWRVHTGEKPYKCNECD RSFSRNSCLALHRRVHTGEKPYKCYECDKVFSR NSCLALHQKTHIGEKPYTCKECGQAFSVRSTLTN HQVIHSDK  3896 A 202 498 MVQSCSAYGCKNRYDKDKPVSFHKFPLTRPSLC KEWEAAVRRKNFKPTKYSSICSEHFIPDCFKREC NNKLLKENAVPTITLCTBPHDKKEDLLEPQEQ 3897 A 2 382 SHGLSRAPHLSAAPAPALASRPCFSSAPCSQGGG GGGPATMIHFILLFSRQGKLRLQKWYITLPDKER KKITREIVQIILSRGHRTSSFVDWKELKLVYKRYA SLYFCCAIENQDNELLTLENVHR SLYFCCAIENQDNELLTLENVHR 3898 A 718 305 SEQEPLLGDTPGSREWDILETBEHYKSRWRSIRIL YLTMFLSSVGFSVVMMSIWPYLQKIDPTADTSFL GWVIASYSLGQMVASPIFGLWSNYTRPKEPLIVS GWVIASYSLGQMVASPIFGLWSNYTRRKEPLIVS LISVAANCLYAYLHIPASHNKYYMLVARGLLGIG GAMAERKGTAKVDFLKKIEKEIQQKWDTERVFE VNASNLEKGYISKGKYFVTFPYPYMNGRLHLGHT FSLSKCEFAVGYQRILKGKCLFPFGLHCTGMPIK ACADKLKREIELY/GCPPDFPDEEEEEEETSVKTE DIIIKDKAKGKKSKAA/AKAGSSKYQWGIMKSLG LSDEEIVKFSEAEHWLDYFNALAIQDLKRMG 3900 A 360 1 VPATSSNVSPSSSESSEPDLSSRSSSSDAPSSSPSVP SPCSLSLSSPESPLLPTLLSSKSPAGSAGPTCGCPS GPGLRATA/PSRLSSSIAAH/SSSAPETSRPAAARE RSPPLHDRESHE 3901 A 193 345 GEWAVPFAPGGQGVSIPHGPEPGQGSGVHIAPRQ                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             | 3895          | A      | 2                                                                        | 1347                                                                     | <u> </u>                                                                                                                                                                                                                                               |
| FRDVAIEFSQDEWKCLNSTQRTLYRDVMLENYR NLVSLDLSRNCVIKELAPQQEGNPJARSIPHSDIGT T*KT*H*RVLLQGNQEKNTRL*LSVER**KKLQQ SDYGPKRKSYL*ERPITR*KRYRKQVY*TSA\*LSF LPHPHELQQFQAEGKIYECNHVEKSVNHGSSVSP PQIISSTIKTHVSNKYGTDFICSSLLTQEQKSCIRE KPYRYIECDKALNHGSHMTVRQVSHSGEKGYKC DLCGKVFSQKSNLARHWRVHTGEKPYKCNECD RSFSRNSCLALHRRVHTGEKPYKCVECDKVFSR NSCLALHQKTHIGEKPYTCKECGQAFSVRSTLTN HQVIHSDK  3896 A 202 498 MVQSCSAYGCKNRYDKDKPVSFHKFPLTRPSLC KEWEAAVRRKNFKPTKYSSICSEHFTPDCFKREC NNKLLKENAVPTIFLCTEPHDKKEDLLEPQEQ 3897 A 2 SHGLSRAPHLSAAPAPALASRPCFSSAPCSQGGG GGGPATMIHFILLFSRQGKLRLQKWYITLPDKER KKITREIVQIILSRGHRTSSFVDWKELKLVYKRYA SLYFCCAIENQDNELLTLENVHR  3898 A 718 305 SEQEPLLGDTPGSREWDILETEEHYKSRWSIRIL YLTMFLSSVGFSVVMMSIWPYLQKIDPTADTSFL GWVIASYSLGQMVASPIFGLWSNYRPKEPLIVSI LISVAANCLYAYLHIPASHNKYYMLVARGLLGIG GAMAERKGTAKVDFLKKIEKEIQQKWDTERVFE VNASNLEKQTSKGKYFVTFPYPYMNGRLHLGHT FSLSKCEFAVGYQRLKGKCCLFPFGLHCTGMPIK ACADKLKREIELY/GCPPDFPDEEEEEEETSVKTE DIIKKDKAKGKSKAA/AKAGSSKYQWGIMKSLG LSDEEIVKFSEAEHWLDYFNALAIQDLKRMG  3900 A 360 1 VPATSSNVSPSSSESSEPDLSSRSSSSDAPSSSPSVP SPCSLSLSSPESPLLPTLLSSKSPAGSAGPTCGCPS GPGLRATA/PSRLSSSIAAH/SSSAPETSRPAAARE RSPPLHDRESHE  3901 A 193 345 GEWAVPPAPGGQGVSIPHGPEPGQGSGVHIAPRQ                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                |               | ļ      |                                                                          |                                                                          |                                                                                                                                                                                                                                                        |
| NLVSLDLSRNCVIKELAPQQEGNP/ARSIPHSDIGT T*KT*H*RVLLQGNQEKNTRL*LSVER**KKLQQ SDYGPKRKSYL*ERPTR*KRYPKQVY*TSA\*LSF LPHPHELQQFQAEGKIYECNHVEKSVNHGSSVSP PQIISSTIKTHVSNKYGTDFICSSLLTQEQKSCIRE KPYRYIECDKALNHGSHMTVRQVSHSGEKGYKC DLCGKVFSQKSNLARHWRVHTGEKPYKCNECD RSFSRNSCLALHRRVHTGEKPYKCYECDKVFSR NSCLALHQKTHIGEKPYTCKECGQAFSVRSTLTN HQVIHSDK  3896 A 202 498 MVQSCSAYGCKNRYDKDKPVSFHKFPLTRPSLC KEWEAAVRRKNFKPTKYSSICSEHFTPDCFKREC NNKLLKENAVPTIFLCTEPHDKKEDLLEPQEQ 3897 A 2 382 SHGLSRAPHLSAAPAPALASRPCFSSAPCSQGGG GGGPATMHFILLFSRQGKLRLQKWYITLPDKER KKITREIVQIILSRGHRTSSFVDWKELKLVYKRYA SLYFCCAIENQDNELLTLENVHR 3898 A 718 305 SEQEPLLGDTPGSREWDILETEEHYKSRWRSIRIL YLTMFLSSVGFSVVMMSIWPYLQKIDPTADTSFL GWVIASYSLGQMVASPIEGLWSNYRPRKEPLIVSI LISVAANCLYAYLHIPASHNKYYMLVARGLLGIG GAMAERKGTAKVDFLKKIEKQQKWDTERVFE VNASNLEKQTSKGKYFVTPYPYMNGRLHLGHT FSLSKCEFAVGYQRLKGKCCLFPFGLHCTGMPIK ACADKLKREIELY/GCPPDFPDEEEEEEETSVKTE DIIKDKAKGKKSKAA/AKAGSSKYQWGIMKSLG LSDEEIVKFSEAEHWLDYFNALAIQDLKRMG  3900 A 360 1 VPATSSNVSPSSSESSEPDLSSRSSSDAPSSSPSVP SPCSLSLSSPESPLLFTLLSSKSPAGSAGPTCGCPS GPGLRATA/PSRLSSSIAAH/SSSAPETSRPAAARE RSPPLHDRESHE 3901 A 193 345 GEWAVPPAPGGQGVSIPHGPEPGQGSGVHIAPRQ                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    |               |        |                                                                          |                                                                          |                                                                                                                                                                                                                                                        |
| T*KT*H*RVLLQGNQEKNTRL*LSVER**KKLQQ SDYGPKRSSYL*ERPTR*KRYRKQVY*TSA\*LSF                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         |               |        |                                                                          | ·                                                                        |                                                                                                                                                                                                                                                        |
| SDYGPKRKSYL*ERPTR*KRYRKQVY*TSAI*LSF LPHPHELQQFQAEGKIYECNHVEKSVNHFGSSVSP PQIISSTIKTHVSNKYGTDFICSSLLTQEQKSCIRE KPYRYIECDKALNHGSHMTVRQVSHSGEKGYKC DLCGKVFSQKSNLARHWRVHTGEKPYKCYECD RSFSRNSCLALHRRVHTGEKPYKCYECDKVFSR NSCLALHQKTHIGEKPYTCKECGQAFSVRSTLTN HQVIHSDK  3896 A 202 498 MVQSCSAYGCKNRYDKDKPVSFHKFPLTRPSLC KEWEAAVRRNFKPTKYSSICSEHFTPDCFKREC NNKLLKENAVPTFILCTEPHDKKEDLLEPQEQ 3897 A 2 382 SHGLSRAPHLSAAPALASSPCFSSAPCSQGGG GGGPATMHFILLFSRQGKLRLQKWYITLPDKER KKITREIVQIILSRGHRTSSFVDWKELKLVYKRYA SLYFCCAIENQDNELLTLENVHR  3898 A 718 305 SEQEPLLGDTPGSREWDILETEEHYKSRWRSIRIL YLTMFLSSVGFSVVMMSIWPYLQKIDPTADTSFL GWVIASYSLGQMVASPIFGLWSNYRPRKEPLIVSI LISVAANCLYAYLHIPASHNKYYMLVARGLLGIG  3899 A 24 718 FRGRPGIPEREGKGNHSFVEVARVIVVDLHSRLG GAMAERKGTAKVDFLKKIEKEIQQKWDTERVFE VNASNLEKQTSKGKYFVTFPYPYMNGRLHLGHT FSLSKCEFAVGYQRLKGKCCLFPFGLHCTGMPIK ACADKLKREIELY/GCPPDFPDEEEEEETSVKTE DIIKDKAKGKKSKAA/AKAGSSKYQWGIMKSLG LSDEEIVKFSEAEHWLDYFNALAIQDLKRMG  3900 A 360 1 VPATSSNVSPSSSESSEPDLSSRSSSSDAPSSSPSVP SPCSLSLSSPESPLLPTLLSSKSPAGSAGPTCGCPS GPGLRATA/PSRLSSSIAAH/SSSAPETSRPAAARE RSPPLHDRESHE  3901 A 193 345 GEWAVPPAPGGQGVSIPHGPEPGQGSGVHIAPRQ                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       | ļ             |        |                                                                          |                                                                          |                                                                                                                                                                                                                                                        |
| LPHPHELQQFQAEGKIYECNHVEKSVNHGSSVSP PQIISSTIKTHYSNKYGTDFICSSLLTQEQKSCIRE KPYRYIECDKALNHGSHMTVRQVSHSGEGYKC DLCGKVFSQKSNLARHWRVHTGEKPYKCNECD RSFSRNSCLALHRRVHTGEKPYKCYECDKVFSR NSCLALHQKTHIGEKPYTCKECGQAFSVRSTLTN HQVIHSDK  3896 A 202 498 MVQSCSAYGCKNRYDKDKPVSFHKFPLTRPSLC KEWEAAVRRKNFKPTKYSSICSEHFTPDCFKREC NNKLLKENAVPTIFLCTEPHDKKEDLLEPQEQ  3897 A 2 382 SHGLSRAPHLSAAPAPALASRPCFSSAPCSQGGG GGGPATMIHFILLFSRQGKLRLQK WYITLPDKER KKITREIVQIILSRGHRTSSFVDWKELKLVYKRYA SLYFCCAIENQDNELLTLENVHR  3898 A 718 305 SEQEPLLGDTFGSREWDILETEEHYKSRWRSIRIL YLTMFLSSVGFSVVMMSIWPYLQKIDPTADTSFL GWVIASYSLGQMVASPIFGLWSNYRPKEPLIVSI LISVAANCLYAYLHIPASHNKYYMLVARGLLGIG GAMAERKGTAKVDFLKKIEKEIQQKWDTERVFE VNASNLEKQTSKGKYFVTFPYPYMNGRLHLGHT FSLSKCEFAVGYQRLKGKCCLFPFGLHCTGMPIK ACADKLKREIELY/GCPPDFPDEEEEEETSVKTE DIIIKDKAKGKKSKAA/AKAGSSKYQWGIMKSLG LSDEEIVKFSEAEHWLDYFNALAIQDLKRMG  3900 A 360 1 VPATSSNVSPSSSESSEPDLSSRSSSSDAPSSSPSVP SPCSLSLSSPESPLLPTLLSSKSPAGSAGPTCGCPS GPGLRATAPSRLSSSIAAH/SSSAPETSRPAAARE RSPPLHDRESHE  3901 A 193 345 GEWAVPPAPGGQGVSIPHGPEPGQGSGVHIAPRQ                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          |               |        |                                                                          |                                                                          |                                                                                                                                                                                                                                                        |
| KPYRYIECDKALNHGSHMTVRQVSHSGEKGYKC DLCGKVFSQKSNLARHWRVHTGEKPYKCNECD RSFSRNSCLALHRKVHTGEKPYKCNECD RSFSRNSCLALHRKVHTGEKPYKCYECDKVFSR NSCLALHQKTHIGEKPYTCKECGQAFSVRSTLTN HQVIHSDK  3896 A 202 498 MVQSCSAYGCKNRYDKDKPVSFHKFPLTRPSLC KEWEAAVRKNFKPTKYSSICSEHFTPDCFKREC NNKLLKENAVPTIFLCTEPHDKKEDLLEPQEQ GGGPATMIHFILLFSRQGKLRLQK WYITLPDKER KKITREIVQIILSRGHRTSSFVDWKELKLVYKRYA SLYFCCAIENQDNELLTLENVHR  3898 A 718 305 SEQEPLLGDTPGSREWDILETEEHYKSRWRSIRIL YLTMFLSSVGFSVVMMSIWPYLQKIDPTADTSFL GWVIASYSLGQMVASPIFGLWSNYRPRKEPLIVSI LISVAANCLYAYLHIPASHNKYYMLVARGLLGIG GAMAERKGTAKVDFLKKIEKEIQQKWDTERVFE VNASNLEKQTSKGKYFVTPYPYMNGRLHLGHT FSLSKCEFAVGYQRLKGKCCLFPFGLHCTGMPIK ACADKLKREIELYJGCPPDFPDEEEEBEETSVKTE DIIKDKAKGKKSKAA/AKAGSSKYQWGIMKSLG LSDEEIVKFSEAEHWLDYFNALAIQDLKRMG  3900 A 360 1 VPATSSNVSPSSSESSEPDLSSRSSSSDAPSSSPSVP SPCSLSLSSPESPLLPTLLSSKSPAGSAGPTCGCPS GPGLRATA/PSRLSSSIAAH/SSSAPETSRPAAARE RSPPLHDRESHE  3901 A 193 345 GEWAVPPAPGGQGVSIPHGPEPGQGSGVHIAPRQ                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     |               |        |                                                                          |                                                                          |                                                                                                                                                                                                                                                        |
| DLCGKVFSQKSNLARHWRVHTGEKPYKCNECD RSFSRNSCLALHRRVHTGEKPYKCYECDKVFSR NSCLALHQKTHIGEKPYTCKECGQAFSVRSTLTN HQVIHSDK  3896 A 202 498 MVQSCSAYGCKNRYDKDKPVSFHKFPLTRPSLC KEWEAAVRRKNFKPTKYSSICSEHFTPDCFKREC NNKLLKENAVPTIFLCTEPHDKKEDLLEPQEQ  3897 A 2 382 SHGLSRAPHLSAAPAPALASRPCFSSAPCSQGGG GGGPATMIHFILLFSRQGKLRLQKWYITLPDKER KKITREIVQIILSRGHRTSSFVDWKELKLVYKRYA SLYFCCAIENQDNELLTLENVHR  3898 A 718 305 SEQEPLLGDTPGSREWDILETEEHYKSRWRSIRIL YLTMFLSSVGFSVVMMSIWPYLQKIDPTADTSFL GWVIASYSLGQMVASPIFGLWSNYRPRKEPLIVSI LISVAANCLYAYLHIPASHNKYYMLVARGLLGIG GAMAERKGTAKVDFLKKIEKEIQQKWDTERVFE VNASNLEKQTSKGKYFVTFPYPYMNGRLHLGHT FSLSKCEFAVGYQRLKGKCCLFPFGLHCTGMPIK ACADKLKREIELY/GCPPDFPDEEEEEEETSVKTE DIIIKDKAKGKKSKAAJAKAGSSKYQWGIMKSLG LSDEEIVKFSEAEHWLDYFNALAIQDLKRMG  3900 A 360 1 VPATSSNVSPSSSESSEPDLSSRSSSDAPSSSPSVP SPCSLSLSSPESPLLPTILLSSKSPAGSAGPTCGCPS GPGLRATA/PSRLSSSIAAH/SSSAPETSRPAAARE RSPPLHDRESHE  3901 A 193 345 GEWAVPPAPGGQGVSIPHGPEPGQGSGVHIAPRQ                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 |               |        |                                                                          |                                                                          |                                                                                                                                                                                                                                                        |
| RSFSRNSCLALHRRVHTGEKPYKCYECDKVFSR NSCLALHQKTHIGEKPYTCKECGQAFSVRSTLTN HQVIHSDK  MVQSCSAYGCKNRYDKDKPVSFHKFPLTRPSLC KEWEAAVRRKNFKPTKYSSICSEHFTPDCFKREC NNKLLKENAVPTIFLCTEPHDKKEDLLEPQEQ  3897 A 2 382 SHGLSRAPHLSAAPAPALASRPCFSSAPCSQGGG GGGPATMIHFILLFSRQGKLRLQKWYITLPDKER KKITREIVQIILSRGHRTSSFVDWKELKLVYKRYA SLYFCCAIENQDNELLTLENVHR  3898 A 718 305 SEQEPLLGDTPGSREWDILETEEHYKSRWRSIRIL YLTMFLSSVGFSVVMMSIWPYLQKIDPTADTSFL GWVIASYSLGQMVASPIFGLWSNYRPRKEPLIVSI LISVAANCLYAYLHIPASHNKYYMLVARGLLGIG GAMAERKGTAKVDFIKKIEKEIQQKWDTERVFE VNASNLEKQTSKGKYFVTFPYPYMNGRLHLGHT FSLSKCEFAVGYQRLKGKCCLFPFGLHCTGMPIK ACADKLKREIELY/GCPPDFPDEEEEEEETSVKTE DIIKDKAKGKKSKAA/AKAGSSKYQWGIMKSLG LSDEEIVKFSEAEHWLDYFNALAIQDLKRMG  3900 A 360 1 VPATSSNVSPSSSSSESPDLSSRSSSSDAPSSSPSVP SPCSLSLSSPESPLLPTLLSSKSPAGSAGPTCGCPS GFGLRATA/PSRLSSSIAAH/SSSAPETSRPAAARE RSPPLHDRESHE  3901 A 193 345 GEWAVPPAPGGQGVSIPHGPEPGQGSGVHIAPRQ                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  |               |        | r.                                                                       |                                                                          |                                                                                                                                                                                                                                                        |
| NSCLALHQKTHIGEKPYTCKECGQAFSVRSTLTN HQVIHSDK  3896 A 202 498 MVQSCSAYGCKNRYDKDKPVSFHKFPLTRPSLC KEWEAAVRRKNFKPTKYSSICSEHFTPDCFKREC NNKLLKENAVPTIFLCTEPHDKKEDLLEPQEQ  3897 A 2 382 SHGLSRAPHLSAAPAPALASRPCFSSAPCSQGGG GGGPATMIHFILLFSRQGKLRLQKWYITLPDKER KKITREIVQIILSRGHRTSSFVDWKELKLVYKRYA SLYFCCAIENQDNELLTLENVHR  3898 A 718 305 SEQEPLLGDTPGSREWDILETEEHYKSRWRSIRIL YLTMFLSSVGFSVVMMSIWPYLQKIDPTADTSFL GWVIASYSLGQMVASPIFGLWSNYRPRKEPLIVSI LISVAANCLYAYLHIPASHNKYYMLVARGLLGIG GAMAERKGTAKVDFLKKIEKEIQQKWDTERVFE VNASNLEKQTSKGKYFVTFPYPYMNGRLHLGHT FSLSKCEFAVGYQRLKGKCCLFPFGLHCTGMPIK ACADKLKREIELY/GCPPDFPDEEEEEEETSVKTE DIIIKDKAKGKKSKAA/AKAGSSKYQWGIMKSLG LSDEEIVKFSEAEHWLDYFNALAIQDLKRMG  3900 A 360 1 VPATSSNVSPSSSESSEPDLSSRSSSSDAPSSSPSVP SPCSLSLSSPESPLLPTILSSKSPAGSAGPTCGCPS GPGLRATA/PSRLSSIAAH/SSSAPETSRPAAARE RSPPLHDRESHE  3901 A 193 345 GEWAVPPAPGGQGVSIPHGPEPGQGSGVHIAPRQ                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     |               |        |                                                                          |                                                                          | _                                                                                                                                                                                                                                                      |
| HQVIHSDK                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       |               |        | 1                                                                        |                                                                          |                                                                                                                                                                                                                                                        |
| 3896   A   202   498   MVQSCSAYGCKNRYDKDKPVSFHKFPLTRPSLC   KEWEAAVRRKNFKPTKYSSICSEHFTPDCFKREC   NNKLLKENAVPTIFLCTEPHDKKEDLLEPQEQ   3897   A   2   382   SHGLSRAPHLSAAPAPALASRPCFSSAPCSQGGG   GGGPATMIHFILLFSRQGKLRLQK WYITLPDKER   KKITREIVQIILSRGHRTSSFVDWKELKLVYKRYA   SLYFCCAIENQDNELLTLENVHR   SEQEPLLGDTPGSREWDILETEEHYKSRWRSIRIL   YLTMFLSSVGFSVVMMSIWPYLQKIDPTADTSFL   GWVIASYSLGQMVASPIFGLWSNYRPRKEPLIVSI   LISVAANCLYAYLHIPASHNKYYMLVARGLLGIG   GAMAERKGTAKVDFLKKIEKEIQQKWDTERVFE   VNASNLEKQTSKGKYFVTFPYPYMNGRLHLGHT   FSLSKCEFAVGYQRLKGKCCLFPFGLHCTGMPIK   ACADKLKREIELY/GCPPDFPDEEEEEETSVKTE   DIIKDKAKGKKSKAA/AKAGSSKYQWGIMKSLG   LSDEEIVKFSEAEHWLDYFNALAIQDLKRMG   SPCSLSLSSPESPLLSSRSSSDAPSSSPSVP   SPCSLSLSSPESPLLSSRSSSDAPSSSPSVP   SPCSLSLSSPESPLLPTLLSSKSPAGSAGPTCGCPS   GPGLRATA/PSRLSSSIAAH/SSSAPETSRPAAARE   RSPPLHDRESHE   3901   A   193   345   GEWAVPPAPGGQGVSIPHGPEPGQGSGVHIAPRQ                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    | 1             |        |                                                                          |                                                                          |                                                                                                                                                                                                                                                        |
| KEWEAAVRKNFKPTKYSSICSEHFTPDCFKREC NNKLLKENAVPTIFLCTEPHDKKEDLLEPQEQ  3897 A 2 382 SHGLSRAPHLSAAPAALASRPCFSSAPCSQGGG GGGPATMIHFILLFSRQGKLRLQK WYITLPDKER KKITREIVQIILSRGHRTSSFVDWKELKLVYKRYA SLYFCCAIENQDNELLTLENVHR 3898 A 718 305 SEQEPLLGDTPGSREWDILETEEHYKSRWRSIRIL YLTMFLSSVGFSVVMMSIWPYLQKIDPTADTSFL GWVIASYSLGQMVASPIFGLWSNYRPRKEPLIVSI LISVAANCLYAYLHIPASHNKYYMLVARGLLGIG GAMAERKGTAKVDFLKKIEKEIQQKWDTERVFE VNASNLEKQTSKGKYFVTFPYPYMNGRLHLGHT FSLSKCEFAVGYQRLKGKCCLFPFGLHCTGMPIK ACADKLKREIELY/GCPPDFPDEEEEEEETSVKTE DIIIKDKAKGKKSKAA/AKAGSSKYQWGIMKSLG LSDEEIVKFSEAEHWLDYFNALAIQDLKRMG  3900 A 360 1 VPATSSNVSPSSSESSEPDLSSRSSSSDAPSSSPSVP SPCSLSLSSPESPLLPTLLSSKSPAGSAGPTCGCPS GPGLRATA/PSRLSSSIAAH/SSSAPETSRPAAARE RSPPLHDRESHE  3901 A 193 345 GEWAVPPAPGGQGVSIPHGPEPGQGSGVHIAPRQ                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    | 3896          | A      | 202                                                                      | 498                                                                      |                                                                                                                                                                                                                                                        |
| 3897 A 2 382 SHGLSRAPHLSAAPAPALASRPCFSSAPCSQGGG GGGPATMIHFILLFSRQGKLRLQKWYITLPDKER KKITREIVQIILSRGHRTSSFVDWKELKLVYKRYA SLYFCCAIE\NQDNELLTLENVHR  3898 A 718 305 SEQEPLLGDTPGSREWDILETEEHYKSRWRSIRIL YLTMFLSSVGFSVVMMSIWPYLQKIDPTADTSFL GWVIASYSLGQMVASPIFGLWSNYRPKEPLIVSI LISVAANCLYAYLHIPASHNKYYMLVARGLLGIG  3899 A 24 718 FRGRPGIPEREGKGNHSFVEVARVIVVDLHSRLG GAMAERKGTAKVDFLKKIEKEIQKWDTERVFE VNASNLEKQTSKGKYFVTFPYPYMNGRLHLGHT FSLSKCEFAVGYQRLKGKCCLFPFGLHCTGMPIK ACADKLKREIELY/GCPPDFPDEEEEEEETSVKTE DIIIKDKAKGKKSKAA/AKAGSSKYQWGIMKSLG LSDEEIVKFSEAEHWLDYFNALAIQDLKRMG  3900 A 360 1 VPATSSNVSPSSSESSEPDLSSRSSSDAPSSSPSVP SPCSLSLSSPESPLLPTLLSSKSPAGSAGPTCGCPS GPGLRATA/PSRLSSSIAAH/SSSAPETSRPAAARE RSPPLHDRESHE  3901 A 193 345 GEWAVPPAPGGQGVSIPHGPEPGQGSGVHIAPRQ                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       |               |        |                                                                          |                                                                          |                                                                                                                                                                                                                                                        |
| GGGPATMIHFILLFSRQGKLRLQKWYITLPDKER KKITREIVQIILSRGHRTSSFVDWKELKLVYKRYA SLYFCCAIE\NQDNELLTLENVHR  3898 A 718 305 SEQEPLLGDTPGSREWDILETEEHYKSRWRSIRIL YLTMFLSSVGFSVVMMSIWPYLQKIDPTADTSFL GWVIASYSLGQMVASPIFGLWSNYRPRKEPLIVSI LISVAANCLYAYLHIPASHNKYYMLVARGLLGIG GAMAERKGTAKVDFLKKIEKEIQQKWDTERVFE VNASNLEKQTSKGKYFVTFPYPYMNGRLHLGHT FSLSKCEFAVGYQRLKGKCCLFPFGLHCTGMPIK ACADKLKREIELY/GCPPDFPDEEEEEEETSVKTE DIIIKDKAKGKKSKAA/AKAGSSKYQWGIMKSLG LSDEEIVKFSEAEHWLDYFNALAIQDLKRMG  3900 A 360 1 VPATSSNVSPSSSESSEPDLSSRSSSSDAPSSSPSVP SPCSLSLSSPESPLLPTLLSSKSPAGSAGPTCGCPS GPGLRATA/PSRLSSSIAAH/SSSAPETSRPAAARE RSPPLHDRESHE  3901 A 193 345 GEWAVPPAPGGQGVSIPHGPEPGQGSGVHIAPRQ                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      |               |        |                                                                          |                                                                          |                                                                                                                                                                                                                                                        |
| KKITREIVQIILSRGHRTSSFVDWKELKLVYKRYA SLYFCCAIENQDNELLTLENVHR  3898 A 718 305 SEQEPLLGDTPGSREWDILETEEHYKSRWRSIRIL YLTMFLSSVGFSVVMMSIWPYLQKIDPTADTSFL GWVIASYSLGQMVASPIFGLWSNYRPRKEPLIVSI LISVAANCLYAYLHIPASHNKYYMLVARGLLGIG  3899 A 24 718 FRGRPGIPEREGKGNHSFVEVARVIVVDLHSRLG GAMAERKGTAKVDFLKKIEKEIQQKWDTERVFE VNASNLEKQTSKGKYFVTFPYPYMNGRLHLGHT FSLSKCEFAVGYQRLKGKCCLFPFGLHCTGMPIK ACADKLKREIELY/GCPPDFPDEEEEEEETSVKTE DIIIKDKAKGKKSKAA/AKAGSSKYQWGIMKSLG LSDEEIVKFSEAEHWLDYFNALAIQDLKRMG  3900 A 360 1 VPATSSNVSPSSSESSEPDLSSRSSSDAPSSSPSVP SPCSLSLSSPESPLLPTLLSSKSPAGSAGPTCGCPS GPGLRATA/PSRLSSSIAAH/SSSAPETSRPAAARE RSPPLHDRESHE  3901 A 193 345 GEWAVPPAPGGQGVSIPHGPEPGQGSGVHIAPRQ                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         | 3897          | Α      | 2                                                                        | 382                                                                      |                                                                                                                                                                                                                                                        |
| SLYFCCAIENQDNELLTLENVHR  3898 A 718 305 SEQEPLLGDTPGSREWDILETEEHYKSRWRSIRIL YLTMFLSSVGFSVVMMSIWPYLQKIDPTADTSFL GWVIASYSLGQMVASPIFGLWSNYRPRKEPLIVSI LISVAANCLYAYLHIPASHNKYYMLVARGLLGIG  3899 A 24 718 FRGRPGIPEREGKGNHSFVEVARVIVVDLHSRLG GAMAERKGTAKVDFLKKIEKEIQQKWDTERVFE VNASNLEKQTSKGKYFVTFPYPYMNGRLHLGHT FSLSKCEFAVGYQRLKGKCCLFPFGLHCTGMPIK ACADKLKREIELY/GCPPDFPDEEEEEEETSVKTE DIIIKDKAKGKKSKAA/AKAGSSKYQWGIMKSLG LSDEEIVKFSEAEHWLDYFNALAIQDLKRMG  3900 A 360 1 VPATSSNVSPSSESSEPDLSSRSSSSDAPSSSPSVP SPCSLSLSSPESPLLPTLLSSKSPAGSAGPTCGCPS GPGLRATA/PSRLSSSIAAH/SSSAPETSRPAAARE RSPPLHDRESHE  3901 A 193 345 GEWAVPPAPGGQGVSIPHGPEPGQGSGVHIAPRQ                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             |               | ,      |                                                                          |                                                                          |                                                                                                                                                                                                                                                        |
| 3898 A 718 305 SEQEPLLGDTPGSREWDILETEEHYKSRWRSIRIL YLTMFLSSVGFSVVMMSIWPYLQKIDPTADTSFL GWVIASYSLGQMVASPIFGLWSNYRPRKEPLIVSI LISVAANCLYAYLHIPASHNKYYMLVARGLLGIG FRGRPGIPEREGKGNHSFVEVARVIVVDLHSRLG GAMAERKGTAKVDFLKKIEKEIQQKWDTERVFE VNASNLEKQTSKGKYFVTFPYPYMNGRLHLGHT FSLSKCEFAVGYQRLKGKCCLFPFGLHCTGMPIK ACADKLKREIELY/GCPPDFPDEEEEEEETSVKTE DIIIKDKAKGKKSKAA/AKAGSSKYQWGIMKSLG LSDEEIVKFSEAEHWLDYFNALAIQDLKRMG  3900 A 360 1 VPATSSNVSPSSSESSEPDLSSRSSSDAPSSSPSVP SPCSLSLSSPESPLLPTLLSSKSPAGSAGPTCGCPS GPGLRATA/PSRLSSSIAAH/SSSAPETSRPAAARE RSPPLHDRESHE  3901 A 193 345 GEWAVPPAPGGQGVSIPHGPEPGQGSGVHIAPRQ                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     | 1             |        |                                                                          |                                                                          |                                                                                                                                                                                                                                                        |
| YLTMFLSSVGFSVVMMSIWPYLQKIDPTADTSFL GWVIASYSLGQMVASPIFGLWSNYRPRKEPLIVSI LISVAANCLYAYLHIPASHNKYYMLVARGLLGIG  S899 A 24 718 FRGRPGIPEREGKGNHSFVEVARVIVVDLHSRLG GAMAERKGTAKVDFLKKIEKEIQQKWDTERVFE VNASNLEKQTSKGKYFVTFPYPYMNGRLHLGHT FSLSKCEFAVGYQRLKGKCCLFPFGLHCTGMPIK ACADKLKREIELY/GCPPDFPDEEEEEEETSVKTE DIIIKDKAKGKKSKAA/AKAGSSKYQWGIMKSLG LSDEEIVKFSEAEHWLDYFNALAIQDLKRMG  S900 A 360 1 VPATSSNVSPSSSESSEPDLSSRSSSDAPSSSPSVP SPCSLSLSSPESPLLPTLLSSKSPAGSAGPTCGCPS GPGLRATA/PSRLSSSIAAH/SSSAPETSRPAAARE RSPPLHDRESHE  3901 A 193 345 GEWAVPPAPGGQGVSIPHGPEPGQGSGVHIAPRQ                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         | 3898          | A      | 718                                                                      | 305                                                                      |                                                                                                                                                                                                                                                        |
| LISVAANCLYAYLHIPASHNKYYMLVARGLLGIG  3899 A 24 718 FRGRPGIPEREGKGNHSFVEVARVIVVDLHSRLG GAMAERKGTAKVDFLKKIEKEIQQKWDTERVFE VNASNLEKQTSKGKYFVTFPYPYMNGRLHLGHT FSLSKCEFAVGYQRLKGKCCLFPFGLHCTGMPIK ACADKLKREIELY/GCPPDFPDEEEEEEETSVKTE DIIIKDKAKGKKSKAA/AKAGSSKYQWGIMKSLG LSDEEIVKFSEAEHWLDYFNALAIQDLKRMG  3900 A 360 1 VPATSSNVSPSSSESSEPDLSSRSSSSDAPSSSPSVP SPCSLSLSSPESPLLPTLLSSKSPAGSAGPTCGCPS GPGLRATA/PSRLSSSIAAH/SSSAPETSRPAAARE RSPPLHDRESHE  3901 A 193 345 GEWAVPPAPGGQGVSIPHGPEPGQGSGVHIAPRQ                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               | ]             |        |                                                                          |                                                                          | YLTMFLSSVGFSVVMMSIWPYLQKIDPTADTSFL                                                                                                                                                                                                                     |
| 3899 A 24 718 FRGRPGIPEREGKGNHSFVEVARVIVVDLHSRLG GAMAERKGTAKVDFLKKIEKEIQQKWDTERVFE VNASNLEKQTSKGKYFVTFPYPYMNGRLHLGHT FSLSKCEFAVGYQRLKGKCCLFPFGLHCTGMPIK ACADKLKREIELY/GCPPDFPDEEEEEEETSVKTE DIIIKDKAKGKKSKAA/AKAGSSKYQWGIMKSLG LSDEEIVKFSEAEHWLDYFNALAIQDLKRMG  3900 A 360 1 VPATSSNVSPSSSESSEPDLSSRSSSSDAPSSSPSVP SPCSLSLSSPESPLLPTLLSSKSPAGSAGPTCGCPS GPGLRATA/PSRLSSSIAAH/SSSAPETSRPAAARE RSPPLHDRESHE  3901 A 193 345 GEWAVPPAPGGQGVSIPHGPEPGQGSGVHIAPRQ                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   | 1             |        |                                                                          |                                                                          | 3                                                                                                                                                                                                                                                      |
| GAMAERKGTAKVDFLKKIEKEIQQKWDTERVFE VNASNLEKQTSKGKYFVTFPYPYMNGRLHLGHT FSLSKCEFAVGYQRLKGKCCLFPFGLHCTGMPIK ACADKLKREIELY/GCPPDFPDEEEEEEETSVKTE DIIIKDKAKGKKSKAA/AKAGSSKYQWGIMKSLG LSDEEIVKFSEAEHWLDYFNALAIQDLKRMG  3900 A 360 1 VPATSSNVSPSSSESSEPDLSSRSSSSDAPSSSPSVP SPCSLSLSSPESPLLPTLLSSKSPAGSAGPTCGCPS GPGLRATA/PSRLSSSIAAH/SSSAPETSRPAAARE RSPPLHDRESHE  3901 A 193 345 GEWAVPPAPGGQGVSIPHGPEPGQGSGVHIAPRQ                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    | 2000          |        | 24                                                                       | 710                                                                      |                                                                                                                                                                                                                                                        |
| VNASNLEKQTSKGKYFVTFPYPYMNGRLHLGHT FSLSKCEFAVGYQRLKGKCCLFPFGLHCTGMPIK ACADKLKREIELY/GCPPDFPDEEEEEEETSVKTE DIIIKDKAKGKKSKAA/AKAGSSKYQWGIMKSLG LSDEEIVKFSEAEHWLDYFNALAIQDLKRMG  VPATSSNVSPSSSESSEPDLSSRSSSSDAPSSSPSVP SPCSLSLSSPESPLLPTLLSSKSPAGSAGPTCGCPS GPGLRATA/PSRLSSSIAAH/SSSAPETSRPAAARE RSPPLHDRESHE  3901 A 193 345 GEWAVPPAPGGQGVSIPHGPEPGQGSGVHIAPRQ                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   | 2699          | A      | 24                                                                       | /18                                                                      |                                                                                                                                                                                                                                                        |
| FSLSKCEFAVGYQRLKGKCCLFPFGLHCTGMPIK ACADKLKREIELY/GCPPDFPDEEEEEEETSVKTE DIIIKDKAKGKKSKAA/AKAGSSKYQWGIMKSLG LSDEEIVKFSEAEHWLDYFNALAIQDLKRMG  VPATSSNVSPSSSESSEPDLSSRSSSSDAPSSSPSVP SPCSLSLSSPESPLLPTLLSSKSPAGSAGPTCGCPS GPGLRATA/PSRLSSSIAAH/SSSAPETSRPAAARE RSPPLHDRESHE  3901 A 193 345 GEWAVPPAPGGQGVSIPHGPEPGQGSGVHIAPRQ                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     | 1             |        |                                                                          |                                                                          |                                                                                                                                                                                                                                                        |
| ACADKLKREIELY/GCPPDFPDEEEEEEETSVKTE DIIIKDKAKGKKSKAA/AKAGSSKYQWGIMKSLG LSDEEIVKFSEAEHWLDYFNALAIQDLKRMG  3900 A 360 1 VPATSSNVSPSSSESSEPDLSSRSSSSDAPSSSPSVP SPCSLSLSSPESPLLPTLLSSKSPAGSAGPTCGCPS GPGLRATA/PSRLSSSIAAH/SSSAPETSRPAAARE RSPPLHDRESHE  3901 A 193 345 GEWAVPPAPGGQGVSIPHGPEPGQGSGVHIAPRQ                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           |               |        |                                                                          |                                                                          |                                                                                                                                                                                                                                                        |
| LSDEEIVKFSEAEHWLDYFNALAIQDLKRMG  3900 A 360 1 VPATSSNVSPSSSESSEPDLSSRSSSSDAPSSSPSVP SPCSLSLSSPESPLLPTLLSSKSPAGSAGPTCGCPS GPGLRATA/PSRLSSSIAAH/SSSAPETSRPAAARE RSPPLHDRESHE  3901 A 193 345 GEWAVPPAPGGQGVSIPHGPEPGQGSGVHIAPRQ                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  |               |        |                                                                          |                                                                          | ACADKLKREIELY/GCPPDFPDEEEEEEETSVKTE                                                                                                                                                                                                                    |
| 3900 A 360 1 VPATSSNVSPSSSESSEPDLSSRSSSDAPSSSPSVP SPCSLSLSSPESPLLPTLLSSKSPAGSAGPTCGCPS GPGLRATA/PSRLSSSIAAH/SSSAPETSRPAAARE RSPPLHDRESHE  3901 A 193 345 GEWAVPPAPGGQGVSIPHGPEPGQGSGVHIAPRQ                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    | [             |        |                                                                          |                                                                          |                                                                                                                                                                                                                                                        |
| SPCSLSLSSPESPLLPTLLSSKSPAGSAGPTCGCPS GPGLRATA/PSRLSSSIAAH/SSSAPETSRPAAARE RSPPLHDRESHE  3901 A 193 345 GEWAVPPAPGGQGVSIPHGPEPGQGSGVHIAPRQ                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      | 2000          |        | 260                                                                      |                                                                          |                                                                                                                                                                                                                                                        |
| GPGLRATA/PSRLSSSIAAH/SSSAPETSRPAAARE RSPPLHDRESHE 3901 A 193 345 GEWAVPPAPGGQGVSIPHGPEPGQGSGVHIAPRQ                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            | 3900          | A.     | 300                                                                      | 1                                                                        |                                                                                                                                                                                                                                                        |
| RSPPLHDRESHE 3901 A 193 345 GEWAVPPAPGGQGVSIPHGPEPGQGSGVHIAPRQ                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 |               |        |                                                                          |                                                                          |                                                                                                                                                                                                                                                        |
| 3901 A 193 345 GEWAVPPAPGGQGVSIPHGPEPGQGSGVHIAPRQ                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              | ,             |        |                                                                          |                                                                          |                                                                                                                                                                                                                                                        |
| GEGSDRTEPLICPKAAP                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              | 3901          | A      | 193                                                                      | 345                                                                      | GEWAVPPAPGGQGVSIPHGPEPGQGSGVHIAPRQ                                                                                                                                                                                                                     |
|                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                | L             |        |                                                                          |                                                                          | GEGSDRTEPLICPKAAP                                                                                                                                                                                                                                      |

| CEO M         | Method | Predicted                                                                                             | Predicted and                                                                                  | Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid,                                                                                                                                                                                                                                                     |
|---------------|--------|-------------------------------------------------------------------------------------------------------|------------------------------------------------------------------------------------------------|-----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| SEQ ID<br>NO: | Method | Predicted beginning nucleotide location corresponding to first amino acid residue of peptide sequence | Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence | E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \possible nucleotide insertion |
| 3902          | A      | 1188                                                                                                  | 1389                                                                                           | NPAARSAAAREGSPALPPPPVS/SSSGLGLLLPLSP<br>PGSHAANPALSPRAPHSHYRPRPRCGPRRRPR                                                                                                                                                                                                                                        |
| 3903          | A      | 63                                                                                                    | 396                                                                                            | NNMRNPHLSSNHYLNLARTETVFARMESVKQRI<br>LAPGKEGLKNFAGKSLGQIYRVLEKKQDTGETIE<br>LTEDGKPL*VPERKAPLCDCTCFGLPRRYIIAIMS<br>GLGFCISFG                                                                                                                                                                                     |
| 3904          | A      | 732                                                                                                   | 1046                                                                                           | AMSECPLILYIHKHIDTYSQSYLFNDLFYPVYSGG<br>RMVTYEHLREVVFGKSEDEHYPLW*VLFGK*YA<br>VAPNALMFIRFM*NCTFVPKLP*VMDLK**LQYK<br>SR                                                                                                                                                                                            |
| 3905          | A      | 46                                                                                                    | 910                                                                                            | QPPPPPPPPPPPPPPPPPPPPARALSHLRLHPDACLFPS PFPLPCSTMPGMMEKGPELLGKNRSANGSAKSP AGGGGSGASSTNGGLHYSEPESGCSSDDEHDVG MRVGAEYQARIPEFDPGATKYTDKDNGGMLVW SPYHSIPDAKLDEYIAIAKEKHGYNVEQALGMLF WHKHNIEKSLADLPNFTPFPDEWTVEDKVLFEQ AFSFHGKSFHRIQQMLPDKTIASLVKYYYSWKK TRSRTSLMDRQARKLANRHNQGDSDDDVEETHP MDGNDSDYDPKKEAKKEGMS      |
| 3906          | A      | 2                                                                                                     | 513                                                                                            | KVCNCCSQELETSFTYVDKNINLEQRNRSSPSAK<br>GHNHPGELGWENPNEWSQEAAISLISEEEDDTSS<br>EATSSGKSIDYGFISAILFLVTGILLVIISYIVPREV<br>TVDPNTVAAREMERLEKESARLGAHLDRCVIAG<br>LCLLTLGGVILSCLLMMSMWKGELYRRNRFAS                                                                                                                      |
| 3907          | A      | 71                                                                                                    | 412                                                                                            | ILIMSNCLQNFLKITSTRLLCSRLCQQLRSKRKFF<br>GTVPISRLHRRVVITGIGLVTPLGVGTHLVWDRLI<br>GGESGIVSLVGEEYKSIPCSVAAYVPRGSDEGQF<br>NEQNFVSKSD                                                                                                                                                                                  |
| 3908          | A      | 77                                                                                                    | 746                                                                                            | LGTLLGWRAPLFSRCLAFHSPFILLNTPKLVKTAE LPPDRNYVLGAHPHGIMCTGFLCNFSTESNGFSQ LFPGLRPWLAVLAGLFYLPVYRDYIMSFGLCPVS RQSLDFILSQPQLGQAVVIMVGGAHEALYSVPGE HCLTLQKRKGFVRLALRHGASLVPVYSFGENDIF RLKAFATGSWQHWCQLTFKKLMGFSPCIFWGR GLFSATSWGLLPFAVPITTV                                                                           |
| 3909          | A      | 1                                                                                                     | 793                                                                                            | FRAAGRPAAAMGDIPVVGLSSWKASPGKVTEAV KEAIDAGYRHFDCAYFYHNEREVGAGIRCKIKE GAVRREDLLIATKLWCTCHKKSLVETACRKSLK ALKLNYLDLYLIHWPMGFKPPHPEWIMSCSELSF CLSHPRVQDLPLDESNMVIPSDTDFLDTWEAME DLVITGLVKNIGVŚNFNHEQLERLLNKPGLRFKP LTNQIECHPYLTQKNLISFCQSRDVSVTAYRPLG GSCEGVDLIDNPVIKRIAKEHGKSPAQILI                                 |
| 3910          | A      | 202                                                                                                   | 705                                                                                            | FFTMHRKKVDNRIRILIENGVAERQRSLFVVVGD<br>RGKDQVVILHHMLSKATVKARPSVLWCYKKEL<br>GFSSHRKKRMRQLQKKIKNGTLNIKQDDPFELFI<br>AATNIRYCYYNETHKILGNTFGMCVLQDFEALTP<br>NLLARTVETVEGGGLVVILLRTMNSLKQLYTVT<br>M                                                                                                                    |
| 3911          | A      | 3                                                                                                     | 723                                                                                            | AGRGARAAGEGGGPFKSRPRPLPSSRSLPAVGGG<br>RYGADKMAAGGAVAAAPECRLLPYALHKWSSF<br>SSTYLPENILVDKPNDQSSRWSSESNYPPQYLILK<br>LERPAIVQNITFGKYEKTHVCNLKKFKVFGGMN<br>EENMTELLSSGLKNDYNKETFTLKHKIDEQMFPC<br>RFIKIVPLLSWGPSFNFSIWYVELSGIDDPDIVQPC                                                                                |

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| SEQ ID | Method | Predicted                                                                                   | Predicted end                                                                    | Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid,                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        |
|--------|--------|---------------------------------------------------------------------------------------------|----------------------------------------------------------------------------------|------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| NO:    |        | beginning nucleotide location corresponding to first amino acid residue of peptide sequence | nucleotide location corresponding to last amino acid residue of peptide sequence | E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \text{\text{\text{\text{-possible}}} nucleotide insertion}                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        |
|        |        |                                                                                             |                                                                                  | LNWYSKYREQEAIRLCLKHFRQHNYTEAFESLQ<br>KKT                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           |
| 3912   | A      |                                                                                             | 461                                                                              | FEKKQLRRPSLFLLGCCSFGIMAPSLWKGLEGIG<br>LFALAHAAFSAAQHRSYMRLTEKEDESLPIDIVL<br>QTLLAFAVTCYGIVHIAGEFKDMDATSELKNKTF<br>DTVRNHPSFYVFNHRGSEYFSGPSDTANSSNQDA<br>LSSNTSLKLRKLESLRR                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          |
| 3913   | A      | 362                                                                                         | 20                                                                               | APGRPEAKVPERSRESGSRRVRGPLLQLRPGRTS<br>RPASGRGRGGAGGSYGKMRKPDSKIVLLGDMN<br>VGKTSLLQRYMERRFPDTVSTVGGAFYLKQWRS<br>YNISIWDTAGEAGAA                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     |
| 3914   | A      |                                                                                             | 7545                                                                             | PGIRVGITSQTGLSSNLQENCSKLAFISSHGTEKQ LQCMPMEGRGRASSSISDLQGKGFEKGTGEKHV PGVGSARHSPQASAGGSPWQRGKAQTRWLGKP DPGRKRRRGSPQEEGGLRVSAAARLLCSGANRC KVLVRQNSTPNTQQPAVHPSTPPSRPLPQAGRCL VAPLRPHPDWVAAKTLAKALRAPGKPWRLAAP SPLGDLGAPGLPGPSTAPRTLSVEEPGVECNQLC LYADVTDPVLCLGQKDPGVEGKHCEKEKISSSK ELKHVHAKSEPSKPARRLSESLHVVDENKNESKI EREHKRRTSTPVIMEGVQEETDTRDVKRQVERSE ICTEEPQKQKSTLKNEKHLKKDDSETPHLKSLLK KEVKSSKEKPEREKTPSEDKLSVKHKYKGDCMH KTGDETELHSSEKGLKVEENIQKQSQTKLSSDD KTERKSKHRNERKLSVLGKDGKPVSEYIIKTDEN VRKENNKKERRLSAEKTKAEHKSRRSSDSKIQK DSLGSKQHGITLQRRSESYSEDKCDMDSTNMDS NLKPEEVVHKEKRRTKSLLEEKLVLKSKSKTQG KQVKVVETELQEGATKQATTPKPDKEKNTEEND SEKQRKSKVEDKPFEETGVEPVLETASSSAHSTQ KDSSHRAKLPLAKEKYKSDKDSTSTRLERKLSD GHKSRSLKHSSKDIKKKDENKSDDKDGKEVDSS HEKARGNSSLMEKKLSRRLCENRRGSLSQEMAK GEEKLAANTLSTPSGSSLQRPKKSGDMTLIPEQEP MEIDSEPGVENVFEVSKTQDNRNNNSHQDIDSEN MKQKTSATVQKDELRTCTADSKATAPAYKPGR GTGVNSNSEKHADHRSTLTKKMHIQSAVSKMNP GEKEPHHRGTTEVNIDSETVHRMLLSAPSENDRV QKNLKNTAAEEHVAQGDATLEHSTNLDSSPSLSS VTVVPLRESYDPDVIPLFDKRTVLEGSTASTSPAD HSALPNQSLTVRESEVLKTSDSKEGGEGFTVDTP AKASITSKRHIPEAHQATLLDGKQGKVIMPLGSK LTGVIVENENITKEGGLVDMAKKENDLNAEPNL KQTIKATVENGKKDGIAVDHVVGLNTEKYAETV KLKHKRSPGKVKDISIDVERRNENSEVDTSAGSG SAPSVLHQRNGQTEDVATGPRRAEKTSVATSTE GKDKDVTLSPVKAGPATTTSSETRQSEVALPCTS IEADEGLIIGTHSRNNPLHVGAEASECTVFAAAEE GGAVVTEGFAESETFLTSTKEGESGECAVAESED RAADLLAVHAVKIEANVNSVVTEEKDDAVTSAG SEEKCDGSLSRDSEIVEGTITTISSEVESDGAVTSAG TEIRAGSISSEEVDGSQGNMMRMGPKKETEGTV TCTGAEGRSDNFVICSVTGAGPREERMVTGAGV VLGDNDAPPGTSASQEGDGSVNDGTEGESAVTS TGITEDGEGPASCTGSEDSSEGFAISSESEENGESA |

| SEQ ID<br>NO: | Method | Predicted beginning nucleotide location corresponding to first amino acid residue of peptide sequence | Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence | Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \-possible nucleotide insertion                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               |
|---------------|--------|-------------------------------------------------------------------------------------------------------|------------------------------------------------------------------------------------------------|------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
|               |        |                                                                                                       |                                                                                                | MDSTVAKEGTNVPLVAAGPCDDEGIVTSTGAKE EDEEGEDVVTSTGRGNEIGHASTCTGLGEESEGV LICESAEGDSQIGTVVEHVEAEAGAAIMNANENN VDSMSGTEKGSKDTDICSSAKGIVESSVTSAVSG KDEVTPVPGGCEGPMTSAASDQSDSQLEKVEDT TISTGLVGGSYDVLVSGEVPECEVAHTSPSEKED EDIITSVENEECDGLMATTASGDITNQNSLAGGK NQGKVLIISTSTTNDYTPQVSAITDVEGGLSDALR TEENMEGTRVTTEEFEAPMPSAVSGDDSQLTASR SEEKDECAMISTSIGEEFELPISSATTIKCAESLQP VAAAVEERATGPVLISTADFEGPMPSAPPEAESP LASTSKEEKDECALISTSIAEECEASVSGVVVESE NERAGTVMEEKDGSGIISTSSVEDCEGPVSSAVP QEEGDPSVTPAEEMGDTAMISTSTSEGCEAVMIG AVLQDEDRLTITRVEDLSDAAIISTSTAECMPISA SIDRHEENQLTADNPEGNGDLSATEVSKHKVPM PSLIAENNCRCPGPVRGGKEPGPVLAVSTEEGHN GPSVHKPSAGQGHPSAVCAEKEEKHGKECPEIGP FAGRGQKESTLHLINAEEKNVLLNSLQKEDKSPE TGTAGGSSTASYSAGRGLEGNANSPAHLRGPEQ TSGQTAKDSSVSSIRYLAAVNTGAIKADDMPPVQ GTVAEHSFLPAEQQGSEDNLKTSTTKCITGQESKI APSHTMIPPATYSVALLAPKCEQDLTIKNDYSGK WTDQASAEKTGDDNSTRKSFPEEGDIMVTVSSE ENVCDIGNEESPLNVLGGLKLKANLKMEAYVPS EEEKNGEILAPPESLCGGKPSGIAELQREPLLVNE SLNVENSGFRTNEEIHSESYNKGEISSGRKDNAE AISGHSVEADPKEVEEEERHMPKRKRKQHYLSSE DEPDDNPDVLDSRIETAQRQCPETEPHATKEENS RDLEELPKTSSETNSTTSRVMEEKDEYSSSETTGE KPEQNDDDTIKSQE |
| 3915          |        | 1                                                                                                     | 7545                                                                                           | PGIRVGITSQTGLSSNLQENCSKLAFISSHGTEKQ LQCMPMEGRGRASSSISDLQGKGFEKGTGEKHV PGVGSARHSPQASAGGSPWQRGKAQTRWLGKP DPGRKRRGSPQEEGGLRVSAAARLLCSGANRC KVLVRQNSTPNTQQPAVHPSTPPSRPLPQAGRCL VAPLRPHPDWVAAKTLAKALRAPGKPWRLAAP SPLGDLGAPGLPGPSTAPRTLSVEEPGVECNQLC LYADVTDPVLCLGQKDPGVEGKHCEKEKISSSK ELKHVHAKSEPSKPARRLSESLHVVDENKNESKI EREHKRRTSTPVIMEGVQEETDTRDVKRQVERSE ICTEEPQKQKSTLKNEKHLKKDDSETPHLKSLLK KEVKSSKEKPEREKTPSEDKLSVKHKYKGDCMH KTGDETELHSSEKGLKVEENIQKQSQQTKLSSDD KTERKSKHRNERKLSVLGKDGKPVSEYIIKTDEN VRKENNKKERRLSAEKTKAEHKSRRSSDSKIQK DSLGSKQHGITLQRRSESYSEDKCDMDSTNMDS NLKPEEVVHKEKRRTKSLLEEKLVLKSKSKTQG KQVKVVETELQEGATKQATTPKPDKEKNTEEND SEKQRKSKVEDKPFEETGVEPVLETASSSAHSTQ KDSSHRAKLPLAKEKYKSDKDSTSTRLERKLSD GHKSRSLKHSSKDIKKKDENKSDDKDGKEVDSS HEKARGNSSLMEKKLSRRLCENRRGSLSQEMAK GEEKLAANTLSTPSGSSLQRPKKSGDMTLIPEQEP MEIDSEPGVENVFEVSKTQDNRNNNSHQDIDSEN                                                                                                                                                                                                                                                  |

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| SEQ ID | Method                                           | Predicted                       | Predicted end | Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid,                                                  |
|--------|--------------------------------------------------|---------------------------------|---------------|--------------------------------------------------------------------------------------------------------------|
| NO:    | Methou                                           | beginning                       | nucleotide    | E-Glutamic Acid, F-Phenylalanine, G-Glycine, H-Histidine,                                                    |
|        |                                                  | nucleotide                      | location      | I=Isoleucine, K=Lysine, L=Leucine, M=Methionine,                                                             |
|        |                                                  | location                        | corresponding | N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, |
|        | 1                                                | corresponding<br>to first amino | to last amino | X=Unknown, *=Stop codon, /=possible nucleotide deletion,                                                     |
|        |                                                  | acid residue of                 | peptide       | >=possible nucleotide insertion                                                                              |
|        |                                                  | peptide<br>sequence             | sequence      |                                                                                                              |
|        | <del>                                     </del> | sequence                        |               | MKQKTSATVQKDELRTCTADSKATAPAYKPGR                                                                             |
|        |                                                  |                                 |               | GTGVNSNSEKHADHRSTLTKKMHIQSAVSKMNP                                                                            |
|        | j                                                |                                 |               | GEKEPIHRGTTEVNIDSETVHRMLLSAPSENDRV                                                                           |
|        |                                                  |                                 |               | QKNLKNTAAEEHVAQGDATLEHSTNLDSSPSLSS                                                                           |
|        | ŀ                                                | 1                               |               | VTVVPLRESYDPDVIPLFDKRTVLEGSTASTSPAD                                                                          |
|        |                                                  |                                 |               | HSALPNQSLTVRESEVLKTSDSKEGGEGFTVDTP                                                                           |
|        | 1                                                |                                 | ļ             | AKASITSKRHIPEAHQATLLDGKQGKVIMPLGSK                                                                           |
|        |                                                  |                                 |               | LTGVIVENENITKEGGLVDMAKKENDLNAEPNL                                                                            |
|        |                                                  |                                 |               | KQTIKATVENGKKDGIAVDHVVGLNTEKYAETV                                                                            |
|        |                                                  |                                 |               | KLKHKRSPGKVKDISIDVERRNENSEVDTSAGSG<br>  SAPSVLHQRNGQTEDVATGPRRAEKTSVATSTE                                    |
|        |                                                  |                                 |               | GKDKDVTLSPVKAGPATTTSSETRQSEVALPCTS                                                                           |
|        |                                                  |                                 |               | IEADEGLIIGTHSRNNPLHVGAEASECTVFAAAEE                                                                          |
|        |                                                  |                                 | ļ             | GGAVVTEGFAESETFLTSTKEGESGECAVAESED                                                                           |
|        |                                                  |                                 |               | RAADLLAVHAVKIEANVNSVVTEEKDDAVTSAG                                                                            |
|        |                                                  | 1                               | ]             | SEEKCDGSLSRDSEIVEGTITFISEVESDGAVTSAG                                                                         |
|        |                                                  | 1                               |               | TEIRAGSISSEEVDGSQGNMMRMGPKKETEGTV                                                                            |
|        |                                                  |                                 |               | TCTGAEGRSDNFVICSVTGAGPREERMVTGAGV                                                                            |
|        |                                                  |                                 |               | VLGDNDAPPGTSASQEGDGSVNDGTEGESAVTS                                                                            |
|        | Ì                                                |                                 | <b>.</b>      | TGITEDGEGPASCTGSEDSSEGFAISSESEENGESA                                                                         |
|        |                                                  |                                 |               | MDSTVAKEGTNVPLVAAGPCDDEGIVTSTGAKE                                                                            |
|        |                                                  |                                 | İ             | EDEEGEDVVTSTGRGNEIGHASTCTGLGEESEGV                                                                           |
|        |                                                  |                                 |               | LICESAEGDSQIGTVVEHVEAEAGAAIMNANENN                                                                           |
|        |                                                  |                                 |               | VDSMSGTEKGSKDTDICSSAKGIVESSVTSAVSG<br>KDEVTPVPGGCEGPMTSAASDQSDSQLEKVEDT                                      |
|        |                                                  |                                 |               | TISTGLVGGSYDVLVSGEVPECEVAHTSPSEKED                                                                           |
|        |                                                  | ĺ                               |               | EDIITSVENEECDGLMATTASGDITNQNSLAGGK                                                                           |
|        |                                                  |                                 | į             | NQGKVLIISTSTTNDYTPQVSAITDVEGGLSDALR                                                                          |
|        |                                                  |                                 | 1             | TEENMEGTRVTTEEFEAPMPSAVSGDDSQLTASR                                                                           |
| 1      |                                                  |                                 | Ì             | SEEKDECAMISTSIGEEFELPISSATTIKCAESLQP                                                                         |
|        |                                                  |                                 | [             | VAAAVEERATGPVLISTADFEGPMPSAPPEAESP                                                                           |
|        |                                                  |                                 | j             | LASTSKEEKDECALISTSIAEECEASVSGVVVESE                                                                          |
|        |                                                  |                                 |               | NERAGTVMEEKDGSGIISTSSVEDCEGPVSSAVP                                                                           |
|        |                                                  | ]                               |               | QEEGDPSVTPAEEMGDTAMISTSTSEGCEAVMIG                                                                           |
|        |                                                  |                                 |               | AVLQDEDRLTITRVEDLSDAAIISTSTAECMPISA                                                                          |
|        |                                                  |                                 | j             | SIDRHEENQLTADNPEGNGDLSATEVSKHKVPM                                                                            |
|        |                                                  | 1                               |               | PSLIAENNCRCPGPVRGGKEPGPVLAVSTEEGHN                                                                           |
|        |                                                  |                                 | }             | GPSVHKPSAGQGHPSAVCAEKEEKHGKECPEIGP<br>FAGRGQKESTLHLINAEEKNVLLNSLQKEDKSPE                                     |
|        |                                                  | 1                               |               | TGTAGGSSTASYSAGRGLEGNANSPAHLRGPEQ                                                                            |
|        |                                                  | 1                               | 1             | TSGQTAKDSSVSSIRYLAAVNTGAIKADDMPPVQ                                                                           |
|        |                                                  |                                 |               | GTVAEHSFLPAEQQGSEDNLKTSTTKCITGQESKI                                                                          |
|        |                                                  | 1                               |               | APSHTMIPPATYSVALLAPKCEQDLTIKNDYSGK                                                                           |
|        |                                                  |                                 |               | WTDQASAEKTGDDNSTRKSFPEEGDIMVTVSSE                                                                            |
|        |                                                  |                                 |               | ENVCDIGNEESPLNVLGGLKLKANLKMEAYVPS                                                                            |
|        |                                                  |                                 |               | EEEKNGEILAPPESLCGGKPSGIAELQREPLLVNE                                                                          |
|        |                                                  |                                 |               | SLNVENSGFRTNEEIHSESYNKGEISSGRKDNAE                                                                           |
|        |                                                  | ]                               | 1             | AISGHSVEADPKEVEEEERHMPKRKRKQHYLSSE                                                                           |
|        |                                                  |                                 | ]             | DEPDDNPDVLDSRIETAQRQCPETEPHATKEENS                                                                           |
|        |                                                  |                                 |               | RDLEELPKTSSETNSTTSRVMEEKDEYSSSETTGE                                                                          |
| 0015   | <u></u>                                          | <u> </u>                        | 660           | KPEQNDDDTIKSQE                                                                                               |
| 3916   | A                                                | 2                               | 773           | GPFGVLWPSAKPGPVTAVEARPPDASDPEGLRG                                                                            |
|        |                                                  |                                 |               | GSPAPLLAPGPLDPSGRLHPAVSMMSYLKQPPYG                                                                           |
|        |                                                  |                                 |               | MNGLGLAGPAMDLLHPSVGYPATPRKQRRERTT<br>FTRSQLDVLEALFAKTRYPDIFMREEVALKINLPE                                     |
|        | ــــــــــــــــــــــــــــــــــــــ           | <del></del>                     | L             | I IIMADA DEUDI UKIKII DILAMETA VEKIMELE                                                                      |

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| SEQ ID<br>NO: | Method | Predicted beginning nucleotide location corresponding to first amino acid residue of peptide sequence | Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence | Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Scrine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion                                                                                                                                                             |
|---------------|--------|-------------------------------------------------------------------------------------------------------|------------------------------------------------------------------------------------------------|------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
|               |        |                                                                                                       |                                                                                                | SRVQVWFKNRRAKCRQQQQSGSGTKSRPAKKK<br>SSPVRESSGSESSGQFTPPAVSSSASSSSSASSSSA<br>NPAAAAAAGLVVAKLPCPLHIFSLCVFIEENRLV<br>SGSWARDIRSVEETDKSGYR                                                                                                                                                                                                                                                                                                                                                                                                   |
| 3917          | A      | 2                                                                                                     | 776                                                                                            | RNIPGRRFRPPGLRRLLKGPHMPREPRGYRTRVP<br>ALRELVPSSHAGSGASEHCQNNRQGSRQHRASR<br>NVQAGGALAPPRHLCGLCSRLHFLKPDLSVRAA<br>PSRAGASVMALRKELLKSIWYAFTALDVEKSGK<br>VSKSQLRVLSHNLYTVLHIPHDPVALEEHFRDDD<br>DGPVSSQGYMPYLNKYILDKVEEGAFVKEHFDE<br>LCWTLTAKKNYRADSNGNSMLSNQDAFRLWCL<br>FNFLSEDKYPLIMDPDEGEYLLKRYS                                                                                                                                                                                                                                           |
| 3918          | A .    | 10                                                                                                    | 318                                                                                            | WQDLVCLGGSRAQEQKPLQQLWNAILLVAMLL<br>CTGLVVQAQRQASRQSQRELGGQVDLFKRRVV<br>RRLASLKTRRCRLSRAAQGLPDPGAETCAVCLD<br>YFCNKQ                                                                                                                                                                                                                                                                                                                                                                                                                      |
| 3919          | A      | 1                                                                                                     | 204                                                                                            | RVLTAINHTLKENLRKFYKGKKDKPLDLRPKKT<br>RAMRRRLNMHEENLKTKKQHRKERLYPLRKYA<br>AKA                                                                                                                                                                                                                                                                                                                                                                                                                                                             |
| 3920          | A      | 1                                                                                                     | 6.54                                                                                           | RCCRSFVAPLQEKVVFGLFFLGAILCLSFSWLFHT<br>VYCHSEGVSRLFSKLDYSGIALLIMGSFVPWLYY<br>SFYCNPQPCFIYLIVICVLGIAAIIVSQWDMFATPQ<br>YRGVRAGVFLGLGLSGIIPTLHYVISEGFLKAATI<br>GQIGWLMLMASLYITGAALYAARIPERFFPGKCD<br>IWFHSHQLFHIFVVAGAFVHFHGVSNLQEFRFMI<br>GGGCSEEDAL                                                                                                                                                                                                                                                                                       |
| 3921          | A      | 1587                                                                                                  | 452                                                                                            | LERDGCGGEEGGSVRSGAGPDSDPRGASSPPAG HRGTAASPRPVAAPSRTPAPPHTRARASPGLPSG PAWRRVQWFSRVSGQVSTLMKATVLMRQPGRV QEIVGALRKGGGDRLQVISDFDMTLSRFAYNGK RCPSSYNILDNSKIISEECRKELTALLHHYYPIEID PHRTVKEKLPHMVEWWTKAHNLLCQQKIQKFQI AQVVRESNAMLREGYKTFFNTLYHNNIPLFIFSA GIGDILEEIIRQMKVFHPNIHIVSNYMDFNEDGFL QGFKGQLIHTYNKNSSACENCGYFQQLEGKTNV ILLGDSIGDLTMADGVPGVQNILKIGFLNDKVEE RRERYMDSYDIVLEKDETLDVVNGLLQHILCQG VQLEMQGP                                                                                                                                    |
| 3922          | A      | 2                                                                                                     | 164                                                                                            | GKIYQRAFGGHSLKFGKGVQAHGCCCVADRTG<br>HSILHTSYGRERPAPVHLRQDT                                                                                                                                                                                                                                                                                                                                                                                                                                                                               |
| 3923          | A      | 2                                                                                                     | 3258                                                                                           | EHATHAYAKLGTRRRHREVTVFVPTWQLKKNR RVRESHFLTKLHSLKMLSITPSQLENGKKITTYD YRFMVKLAEETDGIIVTNEQIHILMNSSKKLMVK DRLLPFTFAGNLFMVPDDPLGRDGPTLDEFLKKP NRLDTDIGNFLKVWKTLPPSSASVTELSDDADSG PLESLPNMEEVREEKEERQDEEQRQGQGTQKAA EEDDLDSSLASVFRVECPSLSEEILRCLSLHDPPD GALDIDLLPGAASPYLGIPWDGKAPCQQVLAHL AQLTIPSNFTALSFFMGFMDSHRDAIPDYEALVG PLHSLLKQKPDWQWDQEHEEAFLALKRALVSAL CLMAPNSQLPFRLEVTVSHVALTAILHQEHSGRK HPIAYTSKPLLPDEESQGPQSGGDSPYAVAWALK HFSRCIGDTPVVLDLSYASRTTADPEVREGRRVS KAWLIRWSLLVQDKGKRALELALLQGLLGENRL LTPAASMPRFFQVLPPFSDLSTFVCIHMSGYCFYR |

CECAD | Maked | Destined | Destined and | Amine said sequence (A=Algaine C=Custaine D=Acquetic Acid

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|---------------|--------|-------------------------------------------------------------------------------------------------------|------------------------------------------------------------------------------------------------|-------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
|               |        |                                                                                                       |                                                                                                | EDEWCAGFGLYVLSPTSPPVSLSFSCSPYTPTYA HLAAVACGLERFGQSPLPVVFLTHCNWIFSLLWE LLPLWRARGFLSSDGAPLPHPSLLSYIISLTSGLSS LPFIYRTSYRGSLFAVTVDTLAKQGAQGGGQWW SLPKDVPAPTVSPHAMGKRPNLLALQLSDSTLAD IIARLQAGQKLSGSSPFSSAFNSLSLDKESGLLMF KGDKKPRVWVVPTQLRRDLIFSVHDIPLGAHQR PEETYKKLRLLGWWPGMQEHVKDYCRSCLFCIP RNLIGSELKVIESPWPLRSTAPWSNLQIEVVGPVT ISEEGHKHVLIVADPNTRWVEAFPLKPYTHTAVA QVLLQHVFARWGVPVRLEAAQGPQFARHVLVS CGLALGAQVASLSRDLQFPCLTSSGAYWEFKRA LKEFIFLHGKKWAASLPLLHLAFRASSTDATPFK VLTGGESRLTEPLWWEMSSANIEGLKMDVFLLQ LVGELLELHWRVADKASEKAENRRFKRESQEKE WNVGDQVLLLSLPRNGSSAKWVGPFYIGDRLSL SLYRIWGFPTPEKLGCIYPSSLMKAFAKSGTPLSF KVLEQ                                  |
| 3924          | A      |                                                                                                       | 1826                                                                                           | MGSVTVRYFCYGCLFTSATWTVLLFVYFNFSEV TQPLKNVPVKGSGPHGPSPKKFYPRFTRGPSRVL EPQFKANKIDDVIDSRVEDPEEGHLKFSSELGMIF NERDQELRDLGYQKHAFNMLISDRLGYHRDVPD TRNAACKEKFYPPDLPAASVVICFYNEAFSALLR TVHSVIDRTPAHLLHEIILVDDDSDFDDLKGELDE YVQKYLPGKIKVIRNTKREGLIRGRMIGAAHATG EVLVFLDSHCEVNVMWLQPLLAAIREDRHTVGC PVIDIISADTLAYSSSPVVRGGFNWGLHFKWDLV PLSELGRAEGATAPIKSPTMAGGLFAMNRQYFH ELGQYDSGMDIWGGENLEISFRIWMCGGKLFIIP CSRVGHIFRKRRPYGSPEGQDTMTHNSLRLAHV WLDEYKEQYFSLRPDLKTKSYGNISERVELRKKL GCKSFKWYLDNVYPEMQISGSHAKPQQPIFVNR GPKRPKVLQRGRLYHLQTNKCLVAQGRPSQKG GLVVLKACDYSDPNQIWIYNEEHELVLNSLLCLD MSETRSSDPPRLMKCHGSGGSQQWTFGKNNRLY QVSVGQCLRAVDPLGQKGSVAMAICDGSSSQQ WHLEG    |
| 3925          | A      | 5386                                                                                                  | 2897                                                                                           | VRWNSKTECYLSIQTQENFPANLNELVNCIVISSL VTTQRKLKAMSLLGSRNQLARAVLNPNPMDFCT KDLLTTTSERIIAYLRDFNEDQKKAIETAYAMVK HSPSVAKICLIHGPPGTGKSKTIVGLLYRLLTENQ RKGHSDENSNAKIKQNRVLVCAPSNAAVDELM KKIILEFKEKCKDKKNPLGNCGDINLVRLGPEKSI NSEVLKFSLDSQVNHRMKKELPSHVQAMHKRK EFLDYQLDELSRQRALCRGGREIQRQELDENISK VSKERQELASKIKEVQGRPQKTQSIIILESHIICCT LSTSGGLLLESAFRGQGGVPFSCVIVDEAGQSCEI ETLTPLIHRCNKLILVGDPKQLPPTVISMKAQEYG YDQSMMARFCRLLEENVEHNMISRLPILQLTVQ YRMHPDICLFPSNYVYNRNLKTNRQTEAIRCSSD WPFQPYLVFDVGDGSERRDNDSYINVQEIKLVM EIIKLIKDKRKDVSFRNIGIITHYKAQKTMIQKDL DKEFDRKGPAEVDTVDAFQGRQKDCVIVTCVRA NSIQGSIGFLASLQRLNVTITRAKYSLFILGHLRTL MENQHWNQLIQDAQKRGAIIKTCDKNYRHDAV |

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|---------------|--------|-------------------------------------------------------------------------------------------------------|------------------------------------------------------------------------------------------------|--------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
|               |        |                                                                                                       |                                                                                                | KILKLKPVLQRSLTHPPTIAPEGSRPQGGLPSSKL<br>DSGFAKTSVAASLYHTPSDSKEITLTVTSKDPERP<br>PVHDQLQDPRLLKRMGIEVKGGIFLWDPQPSSPQ<br>HPGATPPTGEPGFPVVHQDLSHVQQPAAVVAAL<br>SSHKPPVRGEPPAASPEASTCQSKCDDPEEELCH<br>RREARAFSEGEQEKCGSETHHTRRNSRWDKRTL<br>EQEDSSSKKRKLL                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            |
| 3926          | A      | 99                                                                                                    | 284                                                                                            | MPREDRATWKSNYFLKIIQLLDDYPKRFIVGANN<br>VGSKQMQQIRMSLRGKAVVLMGKNTMMR                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           |
| 3927          | A      | 542                                                                                                   | 2                                                                                              | AHLLMLNLAL\TDLL\YLTSLPFLIHYYASGENWI<br>FGDFMCKFIRFSFHFNLYSSILFLTCFSIFRYCVIIH<br>PMSCFSIHKTRCAVVACAVVWIISLVAVIPMTFLI<br>TSTNRTNRSACLDLTSSDELNTIKWYNLILTA\LL<br>CLPLVIVTLCYTTIIHTLTHGHAN\DSCLKQKARR<br>LTILLL                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  |
| 3928          | A      | 1                                                                                                     | 1516                                                                                           | GEEAVGGGAEGGFGVGAQGRAGGRGVEAGR MRLSKTLVDMDMADYSAALDPAYTTLEFENVQ VLTMGNDTSPSEGTNLNAPNSLGVSALCAICGDR ATGKHYGASSCDGCKGFFRRSVRKNHMYSCRFS RQCVVDKDKRNQCRYCRLKKCFRAGMKKEAV QNERDRISTRRSSYEDSSLPSINALLQAEVLSRQIT SPVSGINGDIRAKKIASIADVCESMKEQLLVLVE WAKYIPGFCELPLDDQGALLRAHAGEHLLLGAT KRSMVFKDVLLLGNDYIVPRHCPELAEMSRVSIR ILDELVLPFQELQIDDNEYAYLKAIIFFDPDAKGL SDPGKIKRLRSQVQVSLEDYINDRQYDSRGRFGE LLLLLPTLQSITWQMIEQIQFIKLFGMAKIDNLLQ EMLLGGSPSDAPHAHHPLHPHLMQEHMGTNVIV ANTMPTHLSNGQMCEWPRPRGQAATPETPQPSP PGASGSEPYKLLPGAVATIVKPLSAIPQPTITKQE VI                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      |
| 3929          | Α      |                                                                                                       |                                                                                                | RVLSLESPLEKDPRVLGAQSVPRGRALKGLSPLG LDSAFRLFPDPRAGPWNTAVLSSGMEPETALWG PDLQGPEQSPNDAHRGAESENEEESPRQESSGEEI IMGDPAQSPESKDSTEMSLERSSQDPSVPQNPPTP LGHSNPLDHQIPLDPPAPEVVPTPSDWTKACEAS WQWGALTTWNSPPVVPANEPSLRELVQGRPAG AEKPYICNECGKSFSQWSKLLRHQRIHTGERPNT CSECGKSFTQSSHLVQHQRTHTGEKPYKCPDCG KCFSWSSNLVQHQRTHTGEKPYKCTECEKAFTQ STNLIKHQRSHTGEKPYKCGECRRAFYRSSDLIQ HQATHTGEKPYKCPECGKRFGQNHNLLKHQKIH AGEKPYRCTECGKSFIQSSELTQHQRTHTGEKPY ECLECGKSFGHSSTLIKHQRTHLREDPFKCPVCG KTFTLSATLLRHQRTHTGERPYKCPECGKSFSVS SNLINHQRIHRGERPYICADCGKSFIMSSTLIRHQ RIHTGEKPYKCSDCGKSFIRSSHLIQHRRTHTGEK PYKCPECGKSFSQSSNLITHVRTHMDENLFVCSD CGKAFLEAHELEQHRVIHERGKTPARRAQGDSL LGLGDPSLLTPPPGAKPHKCLVCGKGFNDEGIFM QHQRIHIGENPYKNADGLIAHAAPKPPQLRSPRL PFRGNSYPGAAEGRAEAPGQPLKPPEGQEGFSQR RGLLSSKTYICSHCGESFLDRSVLLQHQLTHGNE KPFLFPDYRIGLGEGAGPSPFLSGKPFKCPECKQS FGLSSELLLHQKVHAGGKSSHKSPELGKSSSVLL                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        |

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| 00'A          |        | 1 30 - 32 - 2                                                                                         | Dung! 1 2                                                                                      | LANGUAGE AND                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    |
|---------------|--------|-------------------------------------------------------------------------------------------------------|------------------------------------------------------------------------------------------------|-----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| SEQ ID<br>NO: | Method | Predicted beginning nucleotide location corresponding to first amino acid residue of peptide sequence | Predicted end nucleotide location corresponding to last amino acid residue of peptide sequence | Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=possible nucleotide insertion                                                                                                                                                                                                            |
|               |        |                                                                                                       |                                                                                                | EHLRSPLGARPYRCSDCRASFLDRVALTRHQETH<br>TQEKPPNPEDPPPEAVTLSTDQEGEGETPTPTESS<br>SHGEGQNPKTLVEEKPYLCPECGAGFTEVAALLL<br>HRSCHPGVSL                                                                                                                                                                                                                                                                                                                                                                                                                                                           |
| 3930          | A .    | 513                                                                                                   | 273                                                                                            | KTQETHIYISEHIFFPFLQGFGNLPICMAKTDLSLS<br>HQPDKKGVPSDFILPISDVRASIGAGFIYPLVGTG<br>SRESPLWL                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 |
| 3931          | A      | 16                                                                                                    | 305                                                                                            | KRRDFLSCWPAFTVLGEARGDQVDWSKLYRDT<br>GLVKMSRKPRASSPFSNNHPSTPKRRGRGKHPLI<br>PGPEALSKFPRQPIREKGPVKEVPGTKGSP                                                                                                                                                                                                                                                                                                                                                                                                                                                                                |
| 3932          | A      | 16                                                                                                    | 305                                                                                            | KRRDFLSCWPAFTVLGEARGDQVDWSKLYRDT<br>GLVKMSRKPRASSPFSNNHPSTPKRRGRGKHPLI<br>PGPEALSKFPRQPIREKGPVKEVPGTKGSP                                                                                                                                                                                                                                                                                                                                                                                                                                                                                |
| 3933          | A      | 334                                                                                                   | 1546                                                                                           | STHASEHWDSALQLAKHLAPDQIPFISKEYAIQLE FAGDYVNALAHYEKGITGDNKEHDEACLAGVA QMSIRMGDIRRGVNQALKHPSRVLKRDCGAILE NMKQFSEAAQLYEKGLYYDKAASVYIRSKNWA KVGDLLPHVSSPKIHLQYAKAKEADGRYKEAVV AYENAKQWQSVIRIYLDHLNNPEKAVNIVRETQ SLDGAKMVARFFLQLGDYGSAIQFLVMSKCNNE AFTLAQQHNKMEIYADIIGSEDTTNEDYQSIALY FEGEKRYLQAGKFFLLCGQYSRALKHFLKCPSSE DNVAIEMAIETVGQAKDELLTNQLIDHLLGEND GMPKDAKYLFRLYMALKQYREAAQTAIIIAREE QSAGNYRNAHDVLFSMYAELKSQKIKIPSEMAT NLMILHSYILVKIHVKNGDHMKGARMLIRVANN ISKFPSHIVPILTSTVIECHRAGLKNSAFSFAAML MRPEYRSKIDAKYKKKIEGMVRRPDISEIEEATTP CPFCKFLLPESELL PTRRPILPLTSPKAISVPSPLQGKQHTLVKSCLSVS |
|               |        |                                                                                                       |                                                                                                | GIGGFLVSLSSRMKLQTLAVSVTALKFWSAYVP CQTQDRDALRLTLEQIDLIRRMCASYSELELVTS AKALNDTQKLACLIGVEGGHSLDNSLSILRTFYM LGVRYLTLTHTCNTPWAESSAKGVHSFYNNISGL TDFGEKVVAEMNRLGMMVDLSHVSDAVARRAL EVSQAPVIFSHSAARGVCNSARNVPDDILQLLEE ERWAFVMVSLFHGELIQWQPIRPMCSTVADHFD HIKAVIGSKFIGIGGDYDGAGKYRKKTTCKAPW RTSSRMSS                                                                                                                                                                                                                                                                                             |
| 3935          | A      | 1                                                                                                     | 883                                                                                            | HETTPAVVQSVLLERGWNKFDKQEQNAEDWNL YWRTSSFRMTEHNSVKPWQQLNHHPGTTKLTR KDCLAKHLKHMRRMYGTSLYQFIPLTFVMPNDY TKFVAEYFQERQMLGTKHSYWICKPAELSRGRG ILIFSDFKDFIFDDMYIVQKYISNPLLIGRYKCDLR IYVCVTGFKPLTIYVYQEGLVRFATEKFDLSNLQ NNYAHLTNSSINKSGASYEKIKEVIGHGCKWTLS RFFSYLRSWDVDDLLLWKKIHRMVILTILAIAPS VPFAANCFELFGFDILIDDNEFHRTG                                                                                                                                                                                                                                                                          |
| 3936          | A      | 203                                                                                                   | 441                                                                                            | HLAHSLGPLPKHYQYCVRYLYYQVTKDVIKEFA<br>DDGVKYLELRSTPRRENATGMTKKTYVESILEGI<br>KQSKQENLDIDV                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 |

| SEQ ID NO:   | Position of end of<br>Signal in Amino Acid<br>Sequence | MaxS (MAXIMUM SCORE) | MeanS (Mean Score)                    |
|--------------|--------------------------------------------------------|----------------------|---------------------------------------|
| 1            | 19                                                     | 0.930                | 0.680                                 |
| 2            | 24                                                     | 0.964                | 0.863                                 |
| 3            | 21                                                     | 0.990                | 0.901                                 |
| 4            | 19                                                     | 0.981                | 0.942                                 |
| 5            | 22                                                     | 0.991                | 0.928                                 |
| <del>5</del> | 21                                                     | 0.956                | 0.843                                 |
| 8            | 122                                                    | 0.913                | 0.718                                 |
| 9            | 17                                                     | 0.997                | 0.969                                 |
| 11           | 19                                                     | 0.930                | 0.680                                 |
| 13           | 36                                                     | 0.983                | 0.863                                 |
|              |                                                        | 0.935                | 0.839                                 |
| 14           | 28                                                     | 0.997                | 0.955                                 |
| 15           | 21                                                     |                      | I                                     |
| 16           | 16                                                     | 0.983                | 0.944                                 |
| 17           | 18                                                     | 0.989                | 0.884                                 |
| 19           | 49                                                     | 0.996                | 0.719                                 |
| 20           | 28                                                     | 0.972                | 0.920                                 |
| 21           | 23                                                     | 0.954                | 0.905                                 |
| 22           | 46                                                     | 0.955                | 0.568                                 |
| 23           | 26                                                     | 0.942                | 0.654                                 |
| 24           | 19                                                     | 0.979                | 0.941                                 |
| 25           | 34                                                     | 0.884                | 0.565                                 |
| 26           | 33                                                     | 0.934                | 0.584                                 |
| 27           | 17                                                     | 0.975                | 0.914                                 |
| 28           | 18                                                     | 0.980                | 0.934                                 |
| 29           | 23                                                     | 0.928                | 0.718                                 |
| 30           | 26                                                     | 0.978                | 0.885                                 |
| 32           | 20                                                     | 0.946                | 0.719                                 |
| 33           | 29                                                     | 0.933                | 0.671                                 |
| 35           | 25                                                     | 0.996                | 0.920                                 |
| 36           | 26                                                     | 0.903                | 0.579                                 |
| 40           | 19                                                     | 0.981                | 0.942                                 |
| 47           | 25                                                     | 0.971                | 0.909                                 |
| 53           | 22                                                     | 0.991                | 0.928                                 |
|              | 24                                                     | 0.960                | 0.808                                 |
| 55           |                                                        | 0.986                | 0.967                                 |
| 60           | 19                                                     |                      | · · · · · · · · · · · · · · · · · · · |
| 78           | 22                                                     | 0.913                | 0.718                                 |
| 86           | 20                                                     | 0.883                | 0.555                                 |
| 87           | 24                                                     | 0.982                | 0.889                                 |
| 88           | 17                                                     | 0.997                | 0.969                                 |
| 115          | 19                                                     | 0.930                | 0.680                                 |
| 134          | 36                                                     | 0.983                | 0.863                                 |
| 136          | 17                                                     | 0.913                | 0.696                                 |
| 137          | 19                                                     | 0.958                | 0.905                                 |
| 140          | 28                                                     | 0.935                | 0.839                                 |
| 143          | 32                                                     | 0.914                | 0.740                                 |
| 153          | 21                                                     | 0.997                | 0.955                                 |
| 154          | 25                                                     | 0.913                | 0.583                                 |
| 155          | 29                                                     | 0.972                | 0.857                                 |
| 169          | 30                                                     | 0.977                | 0.817                                 |
| 170          | 30                                                     | 0.977                | 0.819                                 |
| 171          | 30                                                     | 0.977                | 0.819                                 |
| 175          | 47                                                     | 0.926                | 0.606                                 |
|              | 30                                                     | 0.926                | 0.872                                 |
| 176          | ·                                                      |                      | 0.791                                 |
| 177          | 22                                                     | 0.957                |                                       |
| 192          | 43                                                     | 0.930                | 0.678                                 |

| SEQ ID NO: | Position of end f Signal in Amino Acid Sequence | MaxS (MAXIMUM<br>SCORE) | MeanS (Mean Score) |
|------------|-------------------------------------------------|-------------------------|--------------------|
| 195        | 19                                              | 0.956                   | 0.860              |
| 202        | ,21                                             | 0.982                   | 0.871              |
| 203        | 24                                              | 0.957                   | 0.870              |
| 207        | 23                                              | 0.954                   | 0.905              |
| 224        | 46                                              | 0.955                   | 0.568              |
| 225        | 26                                              | 0.942                   | 0.654              |
| 228        | 45                                              | 0.961                   | 0.839              |
| 231        | 28                                              | 0.994                   | 0.937              |
| 232        | 28                                              | 0.993                   | 0.896              |
| 234        | 19                                              | 0.979                   | 0.942              |
| 235        | 19                                              | 0.979                   | 0.941              |
| 238        | 20                                              | 0.987                   | 0.943              |
| 244        | 23                                              | 0.929                   | 0.683              |
| 250        | 34                                              | 0.884                   | 0.565              |
| 256        | 33                                              | 0.934                   | 0.584              |
| 258        | 25                                              | 0.934                   | 0.729              |
| 259        | 22                                              | 0.969                   | 0.871              |
| 264        | 19                                              | 0.952                   | 0.753              |
| 265        | 17                                              | 0.975                   | 0.914              |
| 266        | 17                                              | 0.975                   | 0.914              |
| 271        | 23                                              | 0.974                   | 0.884              |
| 274        | 13                                              | 0.971                   | 0.834              |
| 275        | 18                                              | 0.980                   | 0.934              |
| 278        | 32                                              | 0.958                   | 0.668              |
| 280        | 24                                              | 0.966                   | 0.881              |
| 281        | 24                                              | 0.966                   | 0.881              |
| 286        | 23                                              | 0.928                   | 0.718              |
| 291        | 35                                              | 0.991                   | 0.824              |
| 293        | $\frac{133}{27}$                                | 0.956                   | 0.806              |
| 294        | 23                                              | 0.952                   | 0.827              |
| 301        | 26                                              | 0.978                   | 0.885              |
| 316        | 20                                              | 0.946                   | 0.719              |
| 320        | 28                                              | 0.978                   | 0.726              |
| 327        | 29                                              | 0.933                   | 0.671              |
| 331        | 48                                              | 0.903                   | 0.571              |
| 345        | 25                                              | 0.996                   | 0.920              |
| 349        | 26                                              | 0.903                   | 0.579              |
| 351        | 24                                              | 0.951                   | 0.876              |
| 352        | 18                                              | 0.944                   | 0.716              |
| 353        | 32                                              | 0.992                   | 0.854              |
| 354        | 27                                              | 0.945                   | 0.817              |
| 355        | 16                                              | 0.922                   | 0.716              |
| 356        | 13                                              | 0.959                   | 0.818              |
| 357        | 23                                              | 0.986                   | 0.878              |
| 358        | 19                                              | 0.904                   | 0.671              |
| 359        | 16                                              | 0.988                   | 0.951              |
| 360        | 15                                              | 0.981                   | 0.938              |
| 361        | 18                                              | 0.944                   | 0.716              |
| 362        | 21                                              | 0.984                   | 0.869              |
| 363        | 40                                              | 0.979                   | 0.813              |
| 364        | 18                                              | 0.883                   | 0.693              |
|            | 22                                              | 0.962                   | 0.908              |
| 365        | 22                                              | 0.961                   | 0.827              |
| 366        |                                                 | <u> </u>                | 0.624              |
| 367        | 44                                              | 0.941                   | <u> </u>           |
| 368        | 20                                              | 0.952                   | 0.791              |
| 369        | 22                                              | 0.949                   | 0.840              |
| 370        | 28                                              | 0.957                   | 0.682              |

| SEQ ID NO: | Positi n of end of Signal in Amino Acid Sequence | MaxS (MAXIMUM<br>SCORE) | MeanS (Mean Score) |
|------------|--------------------------------------------------|-------------------------|--------------------|
| 372        | 28                                               | 0.974                   | 0.894              |
| 373        | 19                                               | 0.972                   | 0.947              |
| 374        | 29                                               | 0.968                   | 0.785              |
| 375        | 19                                               | 0.949                   | 0.897              |
| 377        | 23                                               | 0.962                   | 0.910              |
| 378        | 31                                               | 0.974                   | 0.895              |
| 379        | 26                                               | 0.969                   | 0.939              |
| 380        | 27                                               | 0.945                   | 0.817              |
| 383        | 27                                               | 0.945                   | 0.817              |
| 384        | 25                                               | 0.992                   | 0.877              |
| 385        | 32                                               | 0.983                   | 0.825              |
| 386        | 44                                               | 0.924                   | 0.564              |
| 387`       | 26                                               | 0.971                   | 0.894              |
| 388        | 19                                               | 0.989                   | 0.862              |
| 389        | 24                                               | 0.990                   | 0.947              |
| 390        | 34                                               | 0.942                   | 0.635              |
| 391        | 16 .                                             | 0.922                   | 0.716              |
| 394        | 19                                               | 0.987                   | 0.970              |
| 398        | 36                                               | 0.992                   | 0.866              |
| 404        | 13                                               | 0.959                   | 0.818              |
| 417        | 23                                               | 0.986                   | 0.878              |
| 421        | 19                                               | 0.904                   | 0.671              |
| 425        | 28                                               | 0.971                   | 0.717              |
| 431        | 16                                               | 0.988                   | 0.951              |
| 452        | 18                                               | 0.944                   | 0.716              |
| 459        | 21                                               | 0.991                   | 0.902              |
| 468        | 21                                               | 0.984                   | 0.869              |
| 478        | 40                                               | 0.979                   | 0.813              |
| 486        | 18                                               | 0.883                   | 0.693              |
| 499        | 22                                               | 0.962                   | 0.908              |
| 501        | 19                                               | 0.962                   | 0.877              |
| 514        | 44                                               | 0.941                   | 0.624              |
| 529        | 20                                               | 0.952                   | 0.791              |
| 533        | 39                                               | 0.914                   | 0.719              |
| 548        | 28                                               | 0.957                   | 0.682              |
| 561        | 28                                               | 0.974                   | 0.894              |
| 562        | 28                                               | 0.974                   | 0.893              |
| 564        | 18                                               | 0.949                   | 0.806              |
| 576        | 19                                               | 0.972                   | 0.947              |
| 584        | 29                                               | 0.968                   | 0.785              |
| 585        | 28                                               | 0.973                   | 0.810              |
| 591        | 19                                               | 0.949                   | 0.897              |
| 592        | 24                                               | 0.991                   | 0.954              |
| 594        | 20                                               | 0.985                   | 0.959              |
| 595        | 20                                               | 0.985                   | 0.959              |
| 612        | 23                                               | 0.962                   | 0.910              |
| 619        | 31                                               | 0.974                   | 0.895              |
| 621        | 15                                               | 0.959                   | 0.795<br>0.939     |
| 633        | 26                                               | 0.969                   | 0.842              |
| 640        | 20                                               | 0.949                   |                    |
| 645        | 25                                               | 0.911                   | 0.759              |
| 684        | 25                                               | 0.992                   | 0.877              |
| 691        | 32                                               | 0.983                   | 0.825              |
| 698        | 44                                               | 0.924                   | 0.564              |
| 700        | 19                                               | 0.982                   | 0.941              |
| 710        | 26                                               | 0.971                   | 0.894              |
| 714        | 23                                               | 0.965                   | 0.907              |

| ORO ID NO  | D-141                         | Browe CMA VYRATIRA | Moone (Mann Coom)  |
|------------|-------------------------------|--------------------|--------------------|
| SEQ ID NO: | Position of end of            | MaxS (MAXIMUM      | MeanS (Mean Score) |
|            | Signal in Amino Acid Sequence | SCORE)             |                    |
| 718        | 19                            | 0.989              | 0.862              |
| 725        | 21 .                          | 0.976              | 0.851              |
| 728        | 33                            | 0.961              | 0.895              |
| 734        | 25                            | 0.963              | 0.660              |
| 741        | 34                            | 0.942              | 0.635              |
| 744        | 19                            | 0.959              | 0.924              |
| 747        | 16                            | 0.922              | 0.716              |
| 756        | 26                            | 0.973              | 0.864              |
| 767        | 22                            | 0.986              | 0.943              |
| 768        | 27                            | 0.916              | 0.758              |
| 769        | 19                            | 0.987              | 0.970              |
| 770        | 22                            | 0.981              | 0.933              |
| 771        | 34                            | 0.993              | 0.893              |
| 773        | 20                            | 0.968              | 0.939              |
| 774        | 21                            | 0.971              | 0.945              |
| 778        | 22                            | 0.986              | 0.943              |
| 779        | 32                            | 0.973              | 0.846              |
| 781        | 23                            | 0.950              | 0.857              |
| 785        | 27                            | 0.916              | 0.758              |
| 786        | 27                            | 0.916              | 0.758              |
| 788        | 22                            | 0.981              | 0.933              |
| 793        | 22                            | 0.986              | 0.803              |
| 794        | 39                            | 0.892              | 0.654              |
| 797        | 27                            | 0.965              | 0.847              |
| 810        | 22                            | 0.981              | 0.933              |
| 823        | 34                            | 0.993              | 0.893              |
| 825        | 17                            | 0.962              | 0.778              |
| 837        | 20                            | 0.968              | 0.939              |
| 844        | 25                            | 0.984              | 0.951              |
| 845        | 17                            | 0.919              | 0.706              |
| 846        | 21                            | 0.971              | 0.945              |
| 847        | 21                            | 0.971              | 0.945              |
| 890        | 22                            | 0.986              | 0.943              |
| 893        | 24                            | 0.971              | 0.865              |
| 894        | 24                            | 0.971              | 0.865              |
| 896        | 32                            | 0.973              | 0.846              |
| 899        | 31                            | 0.982              | 0.817              |
| 922        | 15                            | 0.882              | 0.706              |
| 924        | 21                            | 0.975              | 0.948              |
| 925        | 21                            | 0.927              | 0.661              |
| 933        | 20                            | 0.967              | 0.906              |
| 960        | 20                            | 0.967              | 0.906              |
| 967        | 38                            | 0.970              | 0.784              |
| 968        | 47                            | 0.970              | 0.557              |
| 972        | 36                            | 0.945              | 0.775              |

# TABLE 8

| SEQ<br>ID<br>NO: | Method | Predicted beginning nucleotide location c rresp nding t first amino acid residue of peptide sequenc | Predicted end nucleotide location corresponding t last amin acid residue f peptide sequence | Amino acid sequence (A=Alanin C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Is leucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Pr line, Q=Glutamine, R=Arginine, S=Serine, T=Thr onine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unkn wn, *=St p c don, /=possible nucleotide deletion, \=possible nucleotide insertion |
|------------------|--------|-----------------------------------------------------------------------------------------------------|---------------------------------------------------------------------------------------------|-----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| 3955             | A      | 235                                                                                                 | 1272                                                                                        | GPREVLAASSLADGSEEQVMAVALVRERDLSFPG<br>VGDAVVNPTRWHLPAQPEMLYEGGEGRMETLK                                                                                                                                                                                                                                                                                                      |

| SEQ Method | Predicted                                                                                 | Predicted end                                                                    | Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         |
|------------|-------------------------------------------------------------------------------------------|----------------------------------------------------------------------------------|-------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| ID<br>NO:  | beginning nucleotide location c rresponding t first amin acid residue of p ptide sequence | nucleotide location corresponding to last amino acid residue of peptide sequence | Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Isoleucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Thre nine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unknown, *=Stop codon, /=possible nucleotide deletion, \=p ssible nucle tide insertion                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        |
|            |                                                                                           |                                                                                  | DKTLQELEELQNDSEAIDQLALESPEVQDLQLERE MALATNRSLAERNLEFQGPLEISRSNLSDRYQELR KLVERCQEQKAKLEKFSSALQPGTLLDLLQVEGM KIEEESEAMAEKFLEGEVPLETFLENFSSMRMLSH LRRVRVEKLQEVVRKPRASQELAGDAPPPRSPPP V/PPSPPGNTPCG*RAAAATISHASLPFALQPIPQPA CGPHCPWSPATGPFPSSVPALLLQRASGPHLPGSP AWTQGCCGLLLVPTEEHAAPPYGFPPPPGPAWPG Y                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              |
| 3956 A     | 821                                                                                       | 385                                                                              | SICADRTERVGIFFYIPAGTTDEADVTHP*EGHSYL<br>SNHAGIQRSSRP/SHYQGE/WHDNCFTADELQLLT<br>YQLCHTYVRCTRSVSIPAPAYYAHLVAFRARYHL<br>VDKEHDSAEGSHVSGQSNGRDPQALAKAVQIHQ<br>DTLRTMYFA                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           |
| 3957 A     | 4621                                                                                      | 240                                                                              | ELISTFKLLLEKKRSEVMKMKKRYEVGLEKLDSA SSQVATMQMELEALHPQLKVASKEVDEMMIMIE KESVEVAKTEKIVKADETIANEQAMASKAIKDEC DADLAGALPILESALAALDTLTAQDITVVKSMKSP PAGVKLVMEAICILKGIKADKIPDPTGSGKKIEDF WGPAKRLLGDMRFLQSLHEYDKDNIPPAYMNIIR KNYIPNPDFVPEKIRNASTAAEGLCKWVIAMDSY DKVAKIVAPKKIKLAAAEGELKIAMDGLRKKQA ALKEVQDKLARLQDTLELNKQKKADLENQVDLC SKKLERAEQLIGGLGGEKTRWSHTALELGQLYIN LTGDILISSGVVAYLGAFTSTYRQNQTKEWTTLCK GRDIPCSDDCSLMGTLGEAVTIRTWNIAGLPSDSF SIDNGIIIMNARR WPLMIDPQSQANKWIKNMEKA NSLYVIKLSEPDYVRTLENCIQFGTPVLLENVGEE LDPILEPLLLKQTFKQGGSTCIRLGDSTIEYAPDFR FYITTKLRNPHYLPETSVKVTLLNFMITPEGMQDQ LLGIVVAQERPDLEEEKQALILQGAENKRQLKEIE DKILEVLSSSEGNILEDETAIKILSSSKALANEISQK QEVAEETEKKIDTTRMGYRPIAIHSSILFFSLADLA NIEPMYQYSLTWFINLFILSIENSEKSEILAKRLQIL KDHFTYSLYVNVCRSLFEKDKLLFSFCLTINLLLH ERAINKAEWRFLLTGGIGLDNPYANPCTWLPQKS WDEICRLDDLPAFKTIRREFMRLKDGWKKVYDSL EPHHEVFPEEWEDKANEFQRMLIIRCLRPDKVIPM LQEFIINRLGRAFIEPPPFDLAKAFGDSNCCAPLIFV LSPGADPMAALLKFADDQGYGGSKLSSLSLGQGQ GPIAMKMLEKAVKEGTWVVLQNCHLATSWMPT LEKVCEELSPESTHPDFRMWLTSYPSPNFPVSVLQ NGVKMTNEAPKGLRANIIRSYLMDPISDPEFFGSC KKPEEFKKLLYGLCFFHALVQERKFGPLWWNIP YEFNETDLRISVQQLHMFLNQYEELPYEALRYMT GECNYGGRVTDDWDRRTLRSILNKFFNPELVENS DYKFDSSGIYFVPPSGDHKSYIEYTKTLPLTPAPEI FGMNANADITKDQSETQLLFDNILLTQSRSAGAG AKSSDEVVNEVASDILGKLPNNFDIEAAMRRYPT TYTQSMNTVLVQEMGRFNKLLKTIRDSCVNIQKA IKGLAVMSTDLEEVVSSILNVKIPEMWMGKSYPS LKPLGSYVNDFLARLKFLQQWYEVGPPPVFWLSG FFFTQAFLTGAQQNYARKYTIPIDLLGFDYEVMED KEYKHPPEDGVFIHGLFLDGASWNRKIKKLAESH |

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| SEQ<br>ID<br>NO: | Method | Predicted beginning nucleotide location corresponding to first amin acid residue of peptide sequence | Predicted end nucleotide location c rresponding to last amino acid residue f peptide sequence | Amino acid sequence (A=Alanine C=Cysteine, D=Aspartic Acid, E=Glutamic Acid, F=Phenylalanine, G=Glycine, H=Histidine, I=Is leucine, K=Lysine, L=Leucine, M=Methionine, N=Asparagine, P=Proline, Q=Glutamine, R=Arginine, S=Serine, T=Threonine, V=Valine, W=Tryptophan, Y=Tyrosine, X=Unkn wn, *=Stop codon, /=possible nucleotide deletion, \=p ssible nucleotide inserti n                    |
|------------------|--------|------------------------------------------------------------------------------------------------------|-----------------------------------------------------------------------------------------------|-------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
|                  |        |                                                                                                      |                                                                                               | SERRGVLSTTGHSTNFVIA\MTLPSDQPKEHWIGR<br>GVALLCQLNS                                                                                                                                                                                                                                                                                                                                               |
| 3958             | A      | 35                                                                                                   | 529                                                                                           | GADMAKSKNHTTHNQSRKWHRNVIKKPLSQRYK<br>SLKGVDPKFLGNMCFTKKHKKKGLKKMQADSA<br>KAVSTCAKAIEALVKPKEVKPKIPKGVSCELN*LA<br>YIAYPKFWTCACACIAKGLRLCQPKAKAQDQTK<br>AQVQIKAQAAAPASVPTQAPKGAQAPTKASG                                                                                                                                                                                                            |
| 3959             | A      | 1883                                                                                                 | 763                                                                                           | LLVLLLRTNLLIASSTRISRATLTCSPPGIPVDPRVR PRVRSHLVMYLGITTGSLHKAVVSGDSSAHLVEEI QLFPDPEPVRNLQLAPTQGAVFVGFSGGVWRVPR ANCSVYESCVDCVLARDPHCAWDPESRTCCLLSA PNLNSWKQDMERGNPEWACASGPMSRSLRPQSR PQIIKEVLAVPNSILELPCPHLSALASYYWSHGPAA VPEASSTVYNGSLLLIVQDGVGGLYQCWATENGF SYPVISYWVDSQDQTLALDPELAGIPREHVKVPLT RVSGGAALAAQQSYWPHFVTVTVLFALVLSGALI ILVASPLRALRARGKVQGCETLRPGEKAPLSREQH LQSPKECRTSASDVDADNNCLGTEVA |
| 3960             | A      | 1                                                                                                    | 481                                                                                           | SYAAPSLFVKSLYWALAFMAVLLAVSGVVIVVLA<br>SRAGARCQQCPPGWVLSEEHCYYFSAEAQAWEA<br>SQAFCSAYHATLPLLSHTQDFLGRYPVSRHSWVG                                                                                                                                                                                                                                                                                   |

#### TABLE 9

AWRGPQGWHWIDEAPLPPQLLPEDGEDNLDINCG ALEEGTLVAANCSTPRPWVCAKGTQ

| SEQ ID NO: | Accession<br>Number | Species                  | Description                                                        | Smith<br>Waterman<br>Score | % Idenity |
|------------|---------------------|--------------------------|--------------------------------------------------------------------|----------------------------|-----------|
| 3937       | Y27700              | Homo sapiens             | Human secreted protein encoded by gene No. 12.                     | 193                        | 25        |
| 3938       | AF093097            | Homo sapiens             | putative RNA-binding protein Q99                                   | 3881                       | 84        |
| 3939       | AB012308            | Anthocidaris crassispina | B2HC                                                               | 4169                       | 74        |
| 3940       | U10248              | Homo sapiens             | ribosomal protein L29                                              | 787                        | 95        |
| 3941       | Y99418              | Homo sapiens             | Human PRO1317<br>(UNQ783) amino acid<br>sequence SEQ ID<br>NO:277. | 4031                       | 100       |
| 3942       | AL023516            | Gallus gallus            | B locus C type Lectin                                              | 198                        | 35        |

5

#### TABLE 10

| SEQ ID<br>NO: | Accession No. | Description                        | Results*                            |
|---------------|---------------|------------------------------------|-------------------------------------|
| 3937          | PR00049       | WILM'S TUMOUR PROTEIN<br>SIGNATURE | PR00049D 0.00 9.168e-11 209-<br>224 |
| 3942          | BL00615       | C-type lectin domain proteins.     | BL00615A 16.68 6.400e-11 37-<br>55  |

<sup>\*</sup> Results Include in order: accession number subtype; raw score; p-value; position of signature in amino acid sequence

| SEQ ID<br>NO: | PFAM Name      | Description                   | P-Value  | PFAM<br>Score |
|---------------|----------------|-------------------------------|----------|---------------|
| 3938          | Piwi           | Piwi domain                   | 2.6e-150 | 512.7         |
| 3940          | Ribosomal L29e | Ribosomal L29e protein family | 2.3e-19  | 77.8          |
| 3941          | Sema           | Sema domain                   | 4e-181   | 615.1         |
| 3942          | lectin_c       | Lectin C-type domain          | 0.086    | -7.1          |

5

## TABLE 12

| SEQ ID NO: | Position of end of<br>Signal in Amino Acid<br>Sequence | MaxS (Maximum Score) | Means (Mean Score) |
|------------|--------------------------------------------------------|----------------------|--------------------|
| 3941       | 31                                                     | 0.985                | 0.926              |
| 3942       | 21                                                     | 0.974                | 0.894              |

## TABLE 13

10

| SEQ ID NO:<br>of full length<br>nucleotide<br>sequence | SEQ ID<br>NO: of full<br>length<br>peptide<br>sequence | SEQ ID NO:<br>of contig<br>nucleotide<br>sequence | SEQ ID NO:<br>of contig<br>peptide<br>sequence | Priority Docket number corresponding SEQ ID NO: in priority application | SEQ ID NO: in<br>USSN 09/496,914 |
|--------------------------------------------------------|--------------------------------------------------------|---------------------------------------------------|------------------------------------------------|-------------------------------------------------------------------------|----------------------------------|
| 3937                                                   | 3943                                                   | 3949                                              | 3955                                           | 787CIP2G_1                                                              | 787_3587                         |
| 3938                                                   | 3944                                                   | 3950                                              | 3956                                           | 787CIP2G_2                                                              | 787_3813                         |
| 3939                                                   | 3945                                                   | 3951                                              | 3957                                           | 787CIP2G_3                                                              | 787_4462                         |
| 3940                                                   | 3946                                                   | 3952                                              | 3958                                           | 787CIP2G_4                                                              | 787_4887                         |
| 3941                                                   | 3947                                                   | 3953                                              | 3959                                           | 787CIP2G_5                                                              | 787_5794                         |
| 3942                                                   | 3948                                                   | 3954                                              | 3960                                           | 787CIP2G_6                                                              | 787_8743                         |

### TABLE 14

| TISSUE ORIGIN          | LIBRARY/<br>RNA SOURCE | HYSEQ LIBRARY<br>NAME | SEQ ID NOS:     |
|------------------------|------------------------|-----------------------|-----------------|
| adult brain            | GIBCO                  | ABD003                | 3940            |
| adult brain            | Clontech               | ABR006                | 3940            |
| adult brain            | Invitrogen             | ABR014                | 3940            |
| cultured preadipocytes | Strategene             | ADP001                | 3937            |
| adult heart            | GIBCO                  | AHR001                | 3940            |
| adult kidney           | GIBCO                  | AKD001                | 3940            |
| adult lung             | GIBCO                  | ALG001                | 3940            |
| young liver            | GIBCO                  | ALV001                | 3940            |
| adult ovary            | Invitrogen             | AOV001                | 3938, 3940-3941 |
| adult spleen           | GIBCO                  | ASP001                | 3940-3941       |
| testis                 | GIBCO                  | ATS001                | 3940            |
| bone marrow            | Clontech               | BMD001                | 3938, 3940      |
| bone marrow            | Clontech               | BMD004                | 3940            |
| adult cervix           | BioChain               | CVX001                | 3940            |
| endothelial cells      | Strategene             | EDT001                | 3940            |
| fetal brain            | Clontech               | FBR006                | 3940            |
| fetal brain            | Invitrogen             | FBT002                | 3940-3941       |
| fetal heart            | Invitrogen             | FHR001                | 3940            |
| fetal kidney           | Clontech               | FKD001                | 3940            |
| fetal kidney           | Clontech               | FKD002                | 3940            |

| TISSUE ORIGIN                             | LIBRARY/<br>RNA SOURCE | HYSEQ LIBRARY<br>NAME | SEQ ID NOS:      |
|-------------------------------------------|------------------------|-----------------------|------------------|
| fetal liver-spleen                        | Columbia<br>University | FLS001                | 3937, 3940       |
| fetal liver-spleen                        | Columbia<br>University | FLS002                | 3938, 3941       |
| fetal liver-spleen                        | Columbia<br>University | FLS003                | 3940 ·           |
| fetal liver                               | Clontech               | FLV004                | 3940             |
| fetal skin                                | Invitrogen             | FSK001                | 3940-3942        |
| fetal spleen                              | BioChain               | FSP001                | 3940             |
| fetal brain                               | GIBCO                  | HFB001                | 3937, 3940-3941  |
| infant brain                              | Columbia<br>University | IB2002                | 3937, 3939, 3941 |
| leukocyte                                 | GIBCO                  | LUC001                | 3940-3941        |
| leukocyte                                 | Clontech               | LUC003                | 3940-3941        |
| melanoma from cell line ATCC<br>#CRL 1424 | Clontech               | MEL004                | 3940             |
| mammary gland                             | Invitrogen             | MMG001                | 3937, 3940-3941  |
| neuronal cells                            | Strategene             | NTU001                | 3937, 3942       |
| prostate                                  | Clontech               | PRT001                | 3938             |
| rectum                                    | Invitrogen             | REC001                | 3940             |
| salivary gland                            | Clontech               | SALs03                | 3941             |
| small intestine                           | Clontech               | SIN001                | 3940             |
| skeletal muscle                           | Clontech               | SKM001                | 3940             |
| spinal cord                               | Clontech               | SPC001                | 3940             |
| thymus                                    | Clontech               | THMc02                | 3938             |
| thyroid gland                             | Clontech               | THR001                | 3942             |
| uterus                                    | Clontech               | UTR001                | 3940             |

WHAT IS CLAIMED IS:

1. An isolated polynucleotide comprising a nucleotide sequence selected from the group consisting of SEQ ID NO:1-984, 1969-2952, 3937-3942 or 3949-3954, a full length protein coding portion of SEQ ID NO:1-984, 1969-2952, 3937-3942 or 3949-3954, a mature protein coding portion of SEQ ID NO: 1-984, 1969-2952, 3937-3942 or 3949-3954, an active domain coding portion of SEQ ID NO: 1-984, 1969-2952, 3937-3942 or 3949-3954, and complementary sequences thereof.

- 2. An isolated polynucleotide encoding a polypeptide with biological activity, wherein said polynucleotide hybridizes to the polynucleotide of claim 1 under stringent hybridization conditions.
- 3. An isolated polynucleotide encoding a polypeptide with biological activity, wherein said polynucleotide has greater than about 90% sequence identity with the polynucleotide of claim 1.
- 4. The polynucleotide of claim 1 wherein said polynucleotide is DNA.
- 5. An isolated polynucleotide of claim 1 wherein said polynucleotide comprises the complementary sequences.
- 6. A vector comprising the polynucleotide of claim 1.
- 7. An expression vector comprising the polynucleotide of claim 1.
- 8. A host cell genetically engineered to comprise the polynucleotide of claim 1.
- 9. A host cell genetically engineered to comprise the polynucleotide of claim 1 operatively associated with a regulatory sequence that modulates expression of the polynucleotide in the host cell.
- 10. An isolated polypeptide, wherein the polypeptide is selected from the group consisting of:
  - (a) a polypeptide encoded by any one of the polynucleotides of claim 1; and
  - (b) a polypeptide encoded by a polynucleotide hybridizing under stringent conditions with any one of SEQ ID NO: 1-984, 1969-2952, 3937-3942 or 3949-3954.

- ... ...... A composition comprising the polypeptide of claim 10 and a carrier.
- An antibody directed against the polypeptide of claim 10. 12.

11.

- A method for detecting the polynucleotide of claim 1 in a sample, comprising: 13.
- a) contacting the sample with a compound that binds to and forms a complex with the polynucleotide of claim 1 for a period sufficient to form the complex; and
- detecting the complex, so that if a complex is detected, the polynucleotide of claim 1 is detected.
- 14. A method for detecting the polynucleotide of claim 1 in a sample, comprising:
- contacting the sample under stringent hybridization conditions with nucleic acid primers that anneal to the polynucleotide of claim 1 under such conditions;
- amplifying a product comprising at least a portion of the polynucleotide of b) claim 1; and
- detecting said product and thereby the polynucleotide of claim 1 in the c) sample.
- 15. The method of claim 14, wherein the polynucleotide is an RNA molecule and the method further comprises reverse transcribing an annealed RNA molecule into a cDNA polynucleotide.
- 16. A method for detecting the polypeptide of claim 10 in a sample, comprising:
- contacting the sample with a compound that binds to and forms a complex with the polypeptide under conditions and for a period sufficient to form the complex; and
- detecting formation of the complex, so that if a complex formation is detected, the polypeptide of claim 10 is detected.
- 17. A method for identifying a compound that binds to the polypeptide of claim 10, comprising:
- a) contacting the compound with the polypeptide of claim 10 under conditions sufficient to form a polypeptide/compound complex; and
- detecting the complex, so that if the polypeptide/compound complex is b) detected, a compound that binds to the polypeptide of claim 10 is identified.

- 18. A method for identifying a compound that binds to the polypeptide of claim 10, comprising:
- a) contacting the compound with the polypeptide of claim 10, in a cell, under conditions sufficient to form a polypeptide/compound complex, wherein the complex drives expression of a reporter gene sequence in the cell; and
- b) detecting the complex by detecting reporter gene sequence expression, so that if the polypeptide/compound complex is detected, a compound that binds to the polypeptide of claim 10 is identified.
- 19. A method of producing the polypeptide of claim 10, comprising,
- a) culturing a host cell comprising a polynucleotide sequence selected fromm the group consisting of SEQ ID NO: 1-984, 1969-2952, 3937-3942 or 3949-3954, a mature protein coding portion of SEQ ID NO: 1-984, 1969-2952, 3937-3942 or 3949-3954, an active domain coding portion of SEQ ID NO: 1-984, 1969-2952, 3937-3942 or 3949-3954, complementary sequences thereof and a polynucleotide sequence hybridizing under stringent conditions to SEQ ID NO: 1-984, 1969-2952, 3937-3942 or 3949-3954, under conditions sufficient to express the polypeptide in said cell; and
  - b) isolating the polypeptide from the cell culture or cells of step (a).
- 20. An isolated polypeptide comprising an amino acid sequence selected from the group consisting of any one of the polypeptides SEQ ID NO: 985-1968, 2953-3936, 3943-3948 or 3955-3960, the mature protein portion thereof, or the active domain thereof.
- 21. The polypeptide of claim 20 wherein the polypeptide is provided on a polypeptide array.
- 22. A collection of polynucleotides, wherein the collection comprising the sequence information of at least one of SEQ ID NO: 1-984, 1969-2952, 3937-3942 or 3949-3954.
- 23. The collection of claim 22, wherein the collection is provided on a nucleic acid array.
- 24. The collection of claim 23, wherein the array detects full-matches to any one of the polynucleotides in the collection.
- 25. The collection of claim 23, wherein the array detects mismatches to any one of the polynucleotides in the collection.

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26. The collection of claim 22, wherein the collection is provided in a computer-readable format.

- 27. A method of treatment comprising administering to a mammalian subject in need thereof a therapeutic amount of a composition comprising a polypeptide of claim 10 or 20 and a pharmaceutically acceptable carrier.
- 28. A method of treatment comprising administering to a mammalian subject in need thereof a therapeutic amount of a composition comprising an antibody that specifically binds to a polypeptide of claim 10 or 20 and a pharmaceutically acceptable carrier.

Pages  $\,485\ \mathrm{to}\ 6221\,$  of this application contain amino acid sequence listings. They can be obtained at the address given below.

Les pages 485 to 6221 de cette demande contiennent des listages des séquences d'acides aminés. Elles peuvent être obtenues à l'adresse indiquée ci-dessous.

World Intellectual Property Organization 34, chemin des Colombettes CH-1211 Genève 20